

US005589136A

United States Patent [19]

Northrup et al.

[11] Patent Number:

5,589,136

[45] Date of Patent:

Dec. 31, 1996

[54] SILICON-BASED SLEEVE DEVICES FOR CHEMICAL REACTIONS

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[21] Appl. No.: **492,678**

[22] Filed: Jun. 20, 1995

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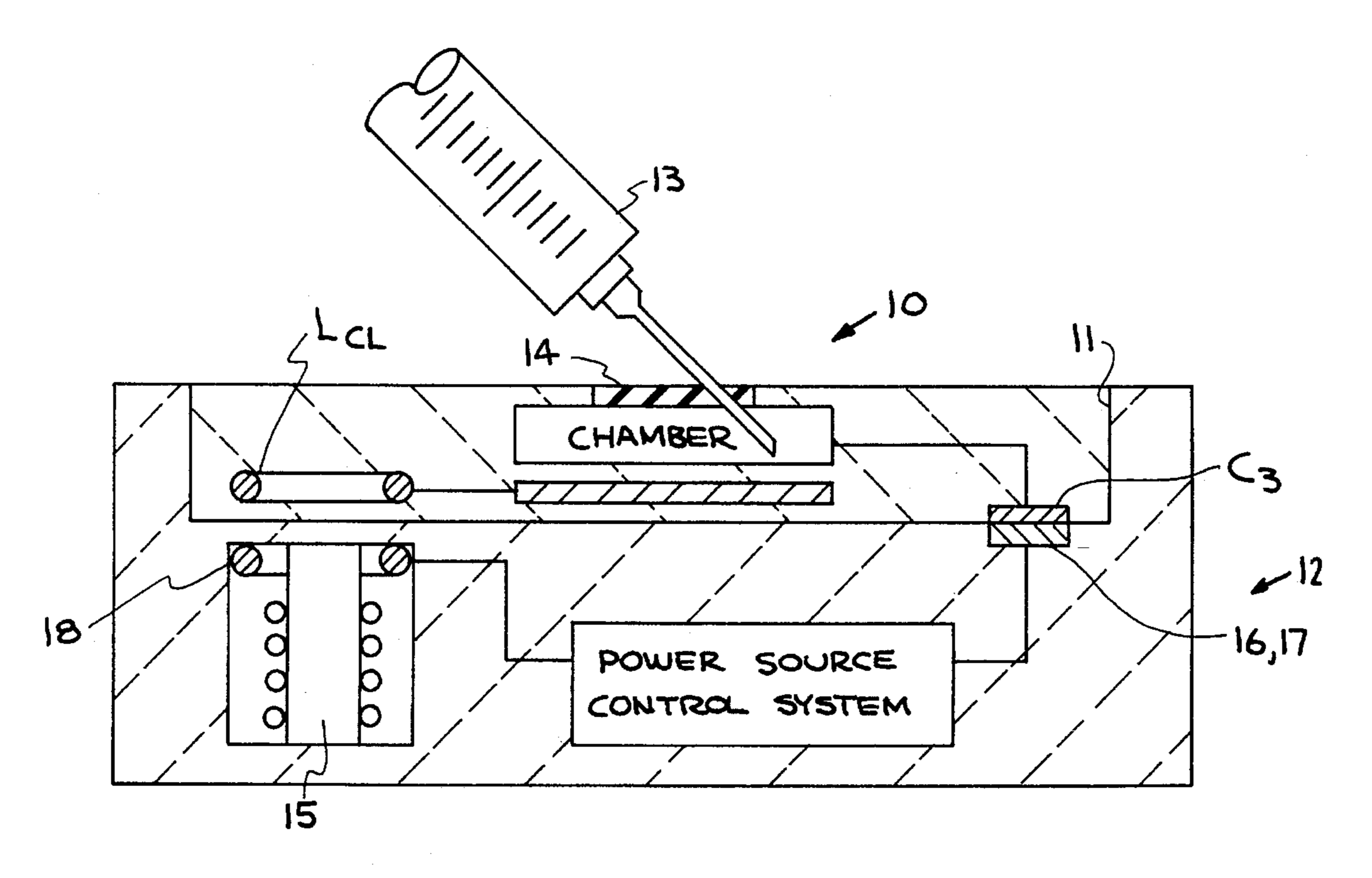
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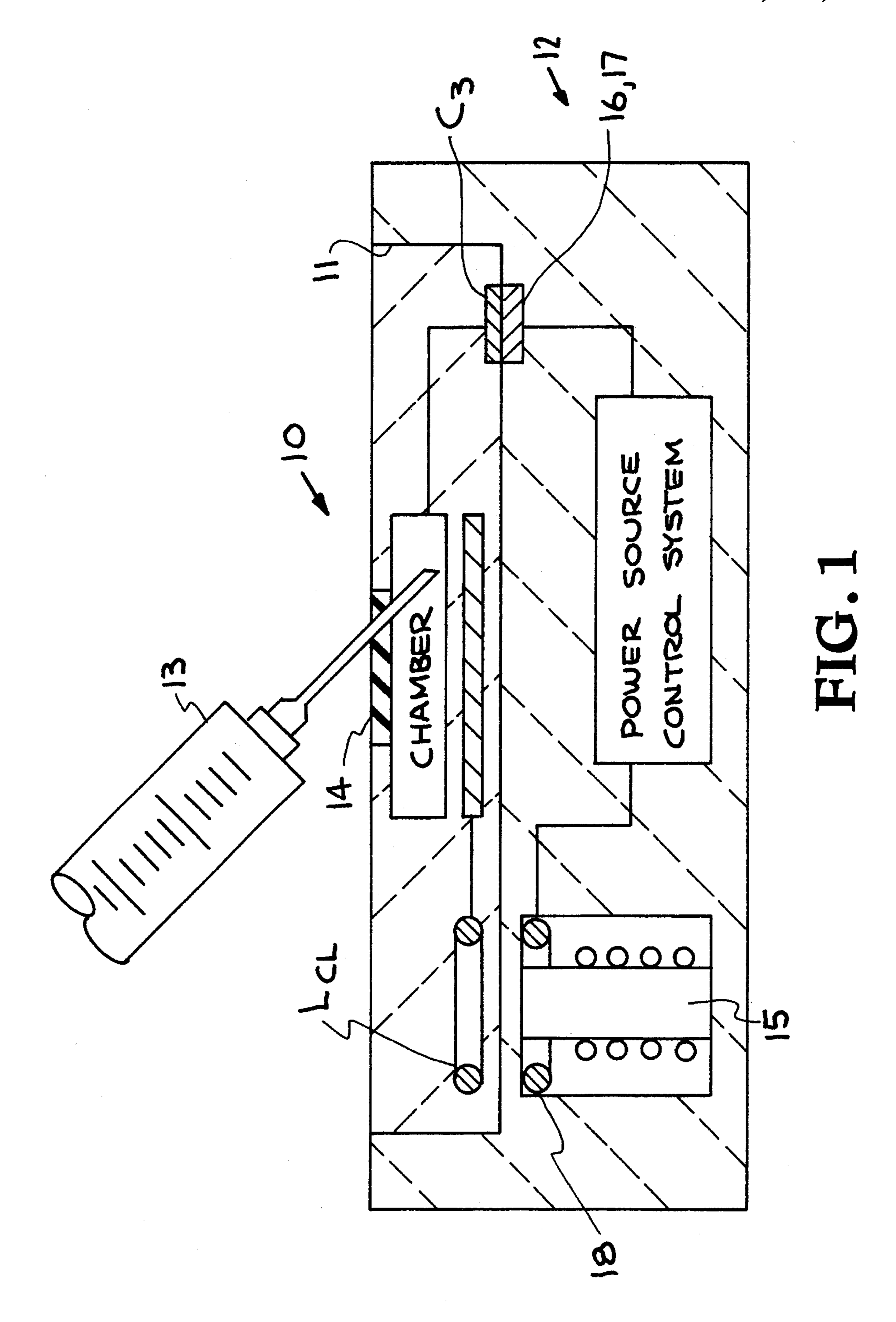
Primary Examiner—Timothy McMahon Attorney, Agent, or Firm—Henry P. Sartorio; L. E. Carnahan

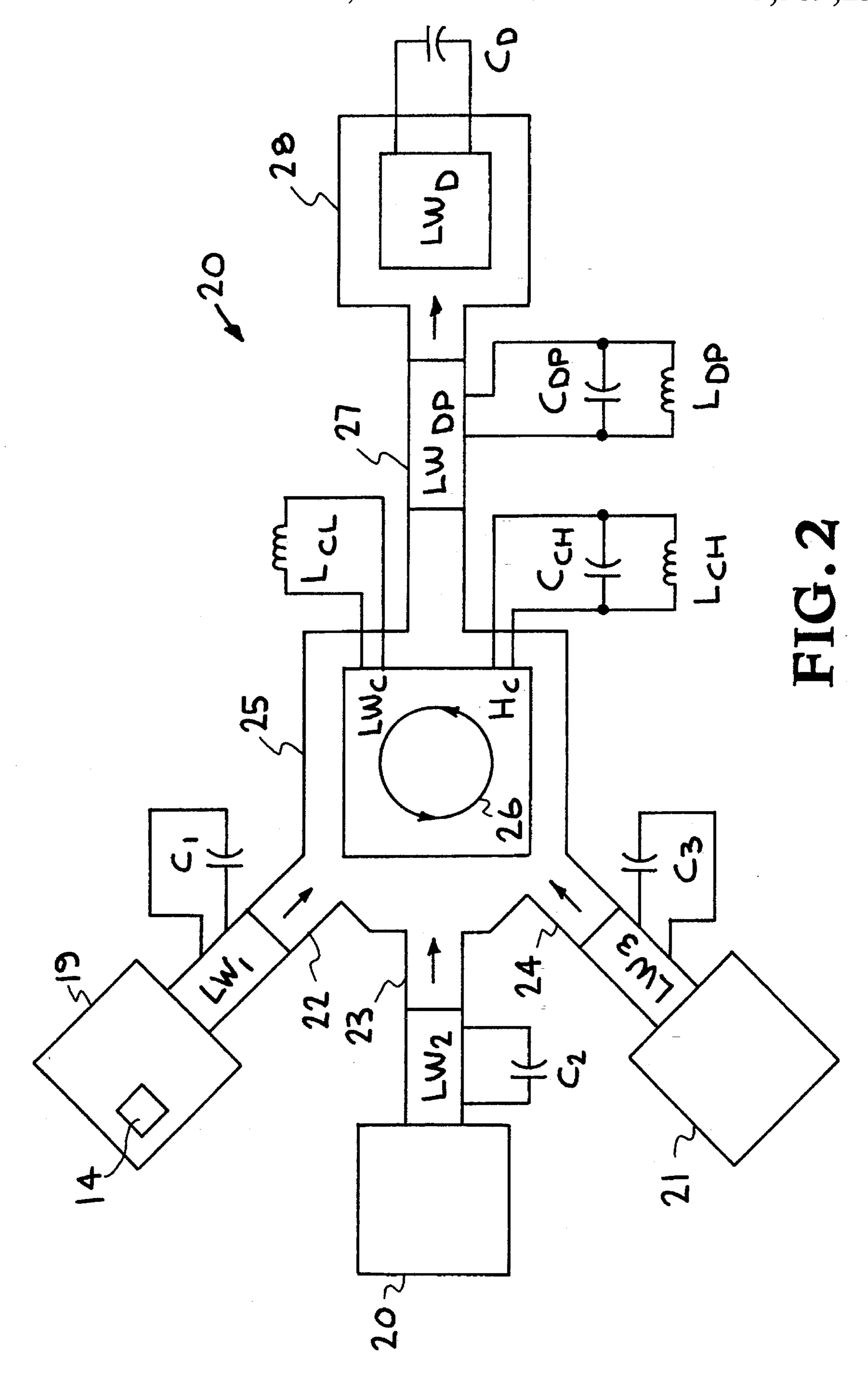
[57] ABSTRACT

A silicon-based sleeve type chemical reaction chamber that combines heaters, such as doped polysilicon for heating, and bulk silicon for convection cooling. The reaction chamber combines a critical ratio of silicon and silicon nitride to the volume of material to be heated (e.g., a liquid) in order to provide uniform heating, yet low power requirements. The reaction chamber will also allow the introduction of a secondary tube (e.g., plastic) into the reaction sleeve that contains the reaction mixture thereby alleviating any potential materials incompatibility issues. The reaction chamber may be utilized in any chemical reaction system for synthesis or processing of organic, inorganic, or biochemical reactions, such as the polymerase chain reaction (PCR) and/or other DNA reactions, such as the ligase chain reaction, which are examples of a synthetic, thermal-cyclingbased reaction. The reaction chamber may also be used in synthesis instruments, particularly those for DNA amplification and synthesis.

22 Claims, 14 Drawing Sheets









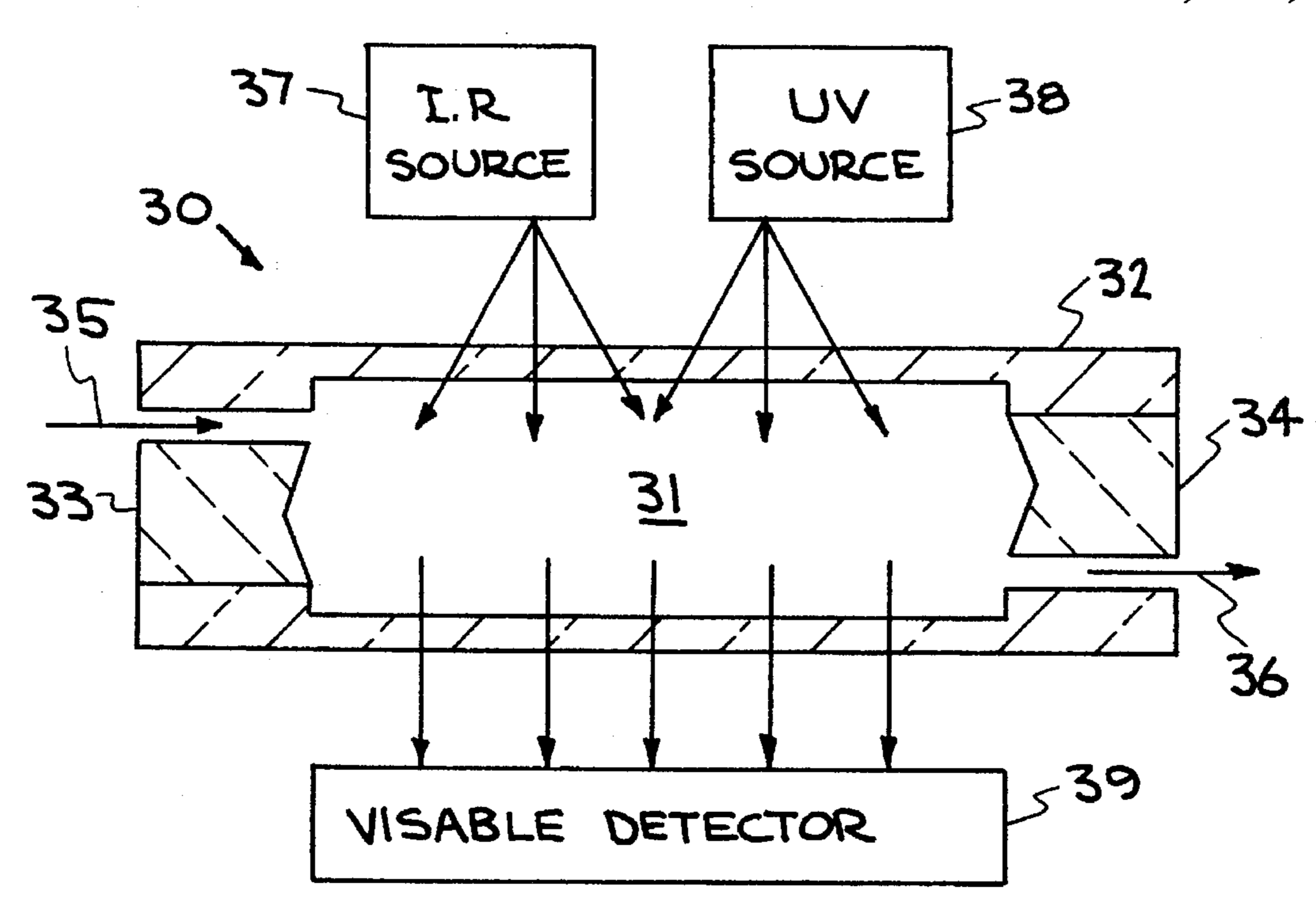
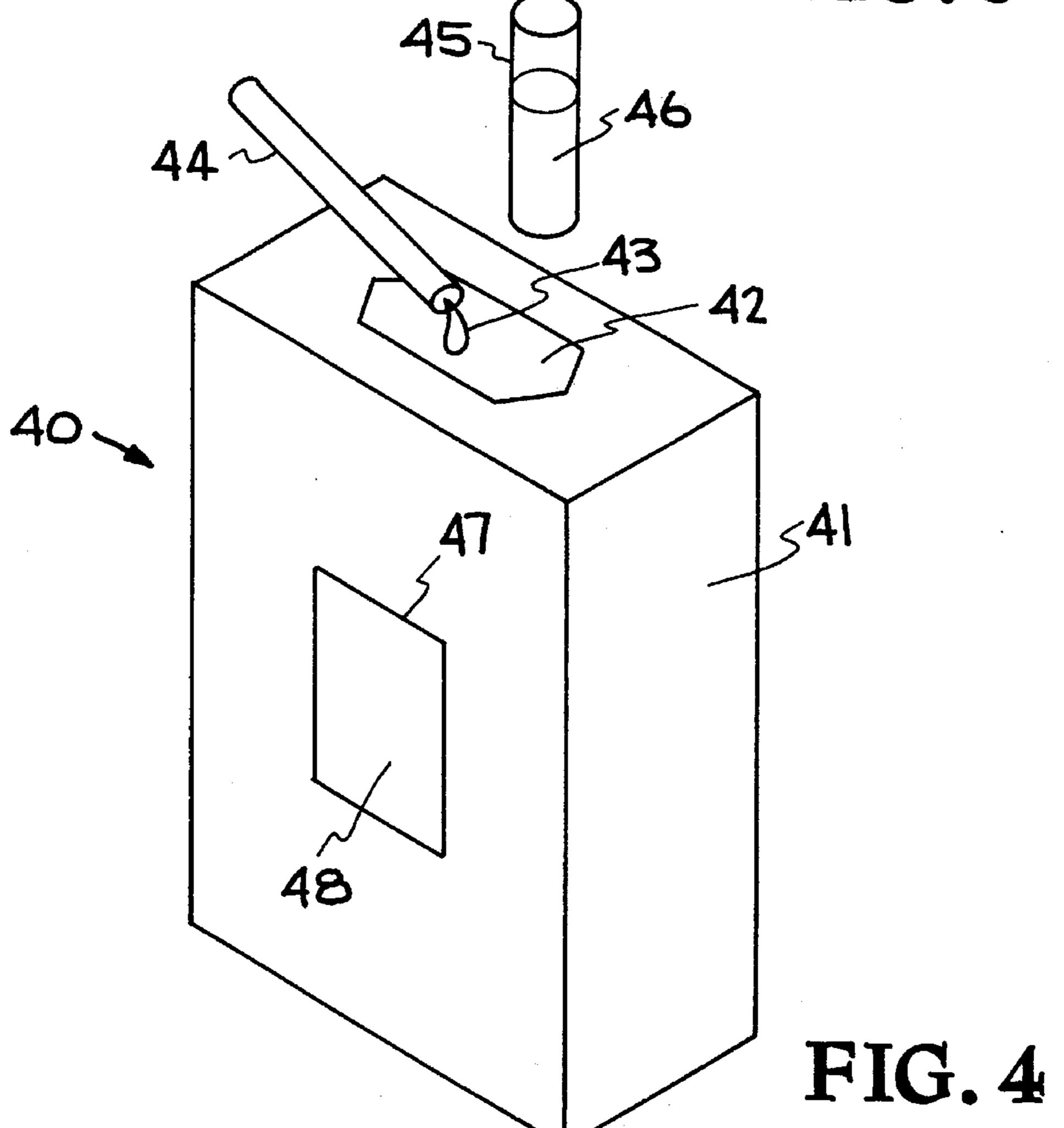
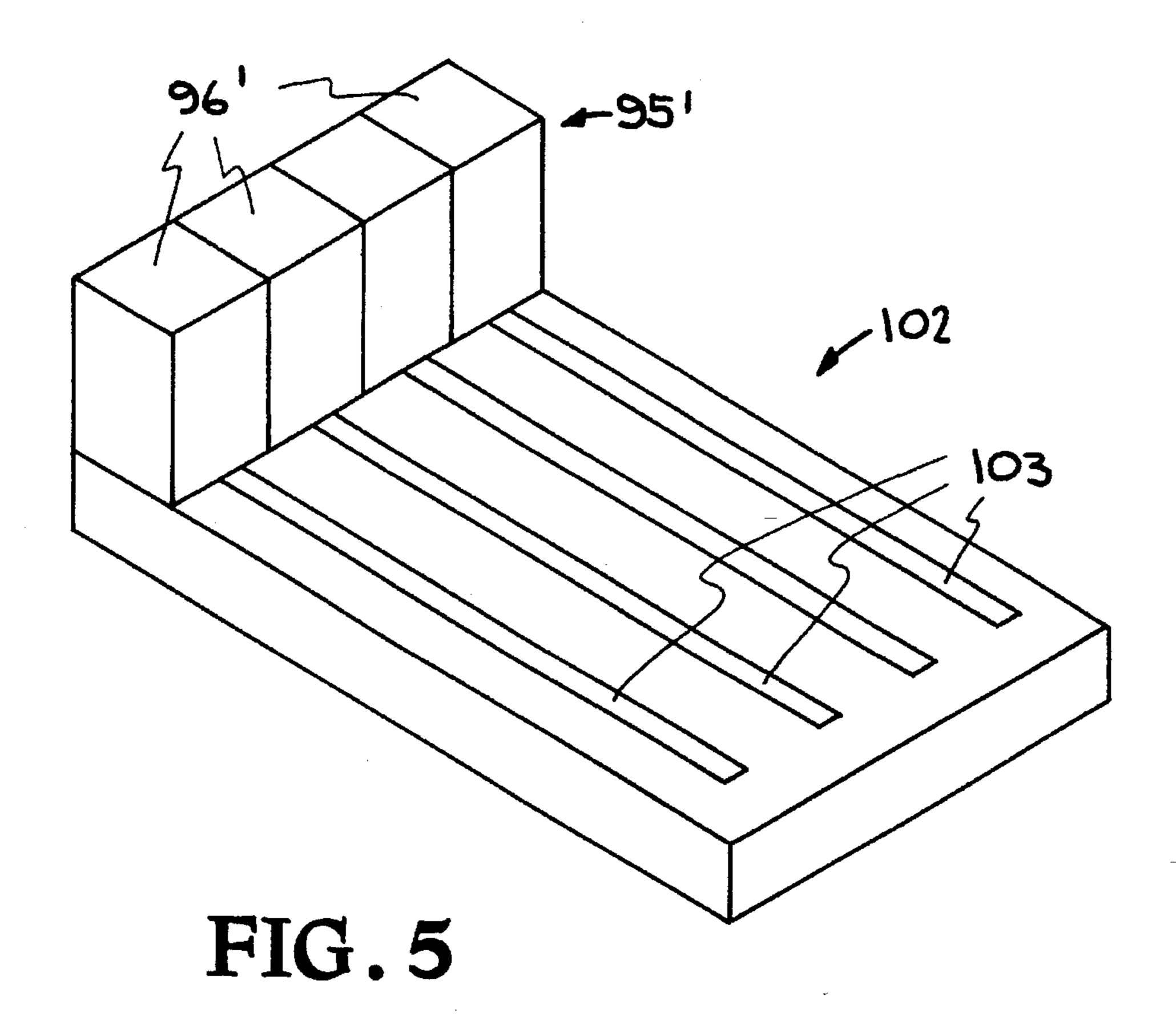
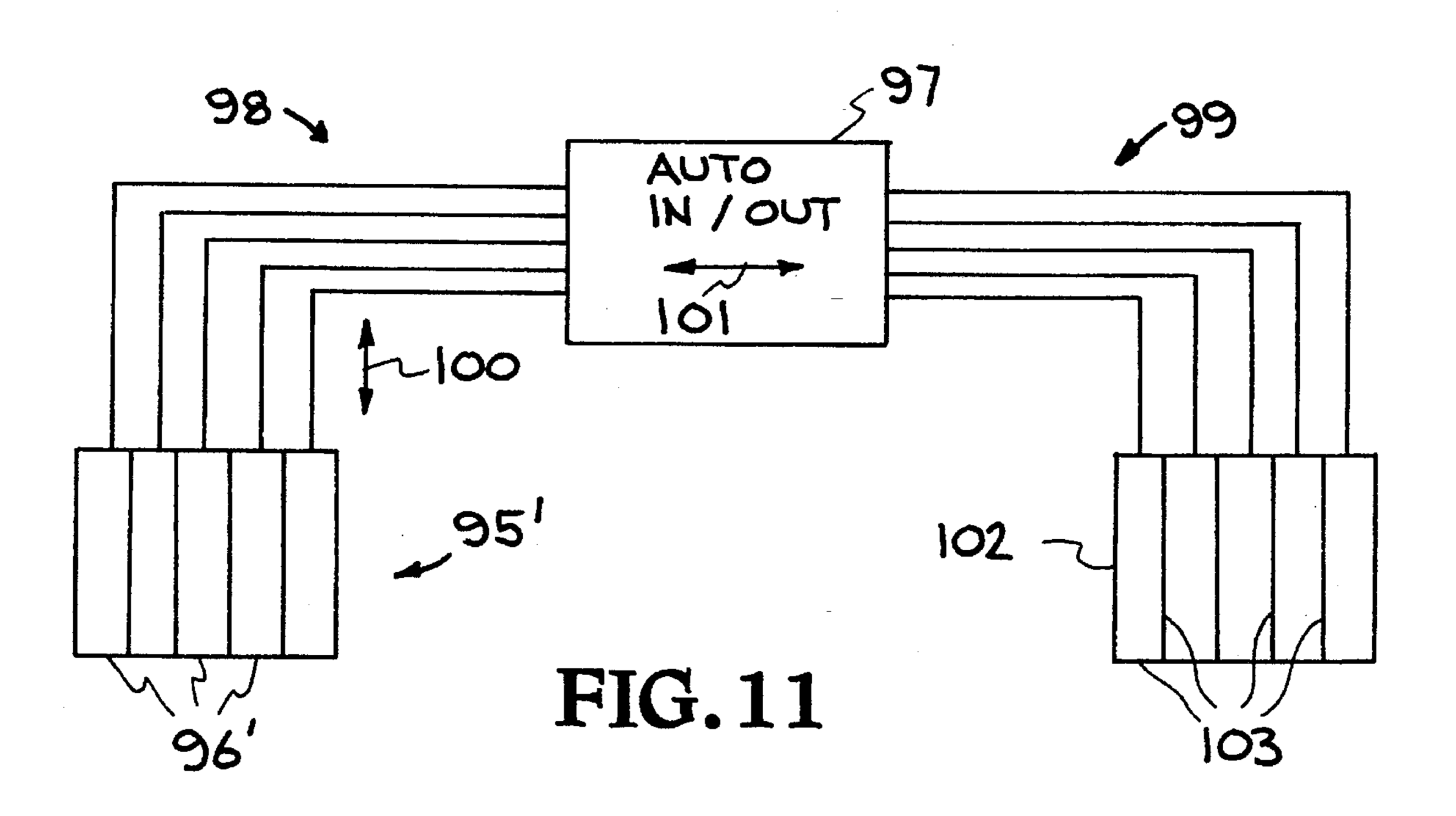
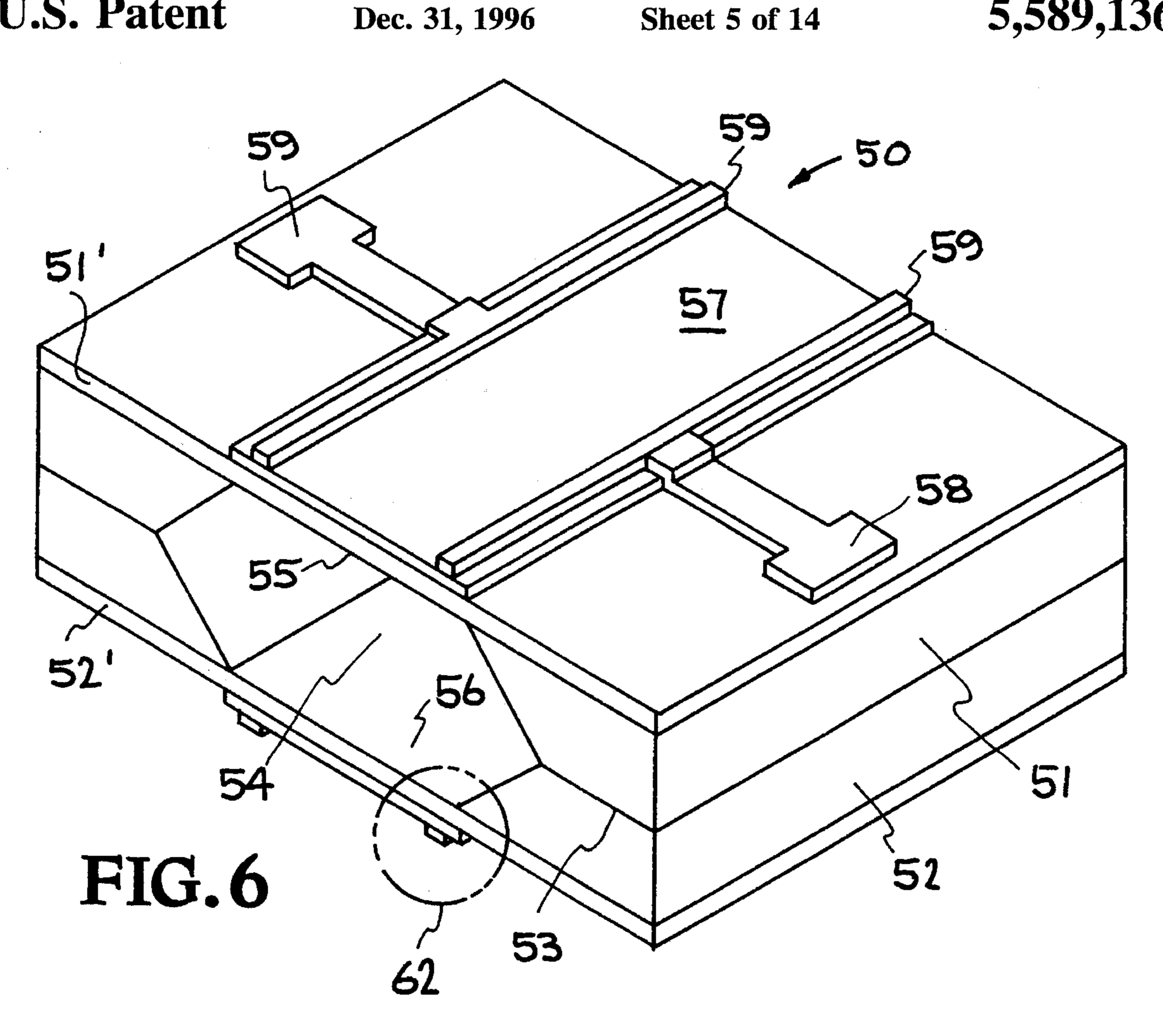


FIG. 3









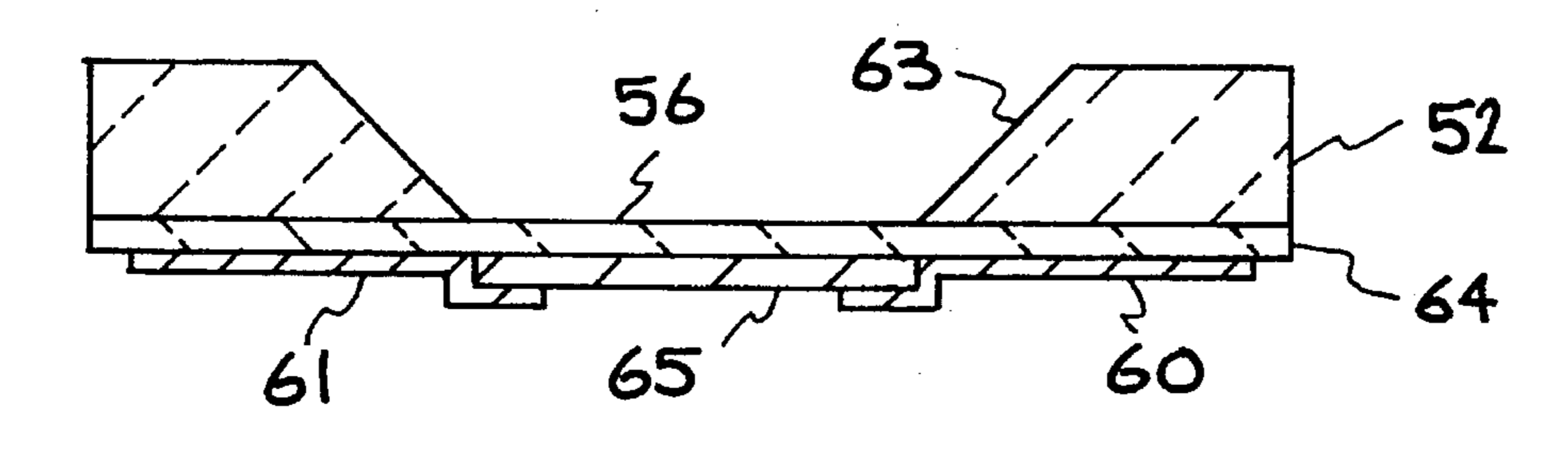
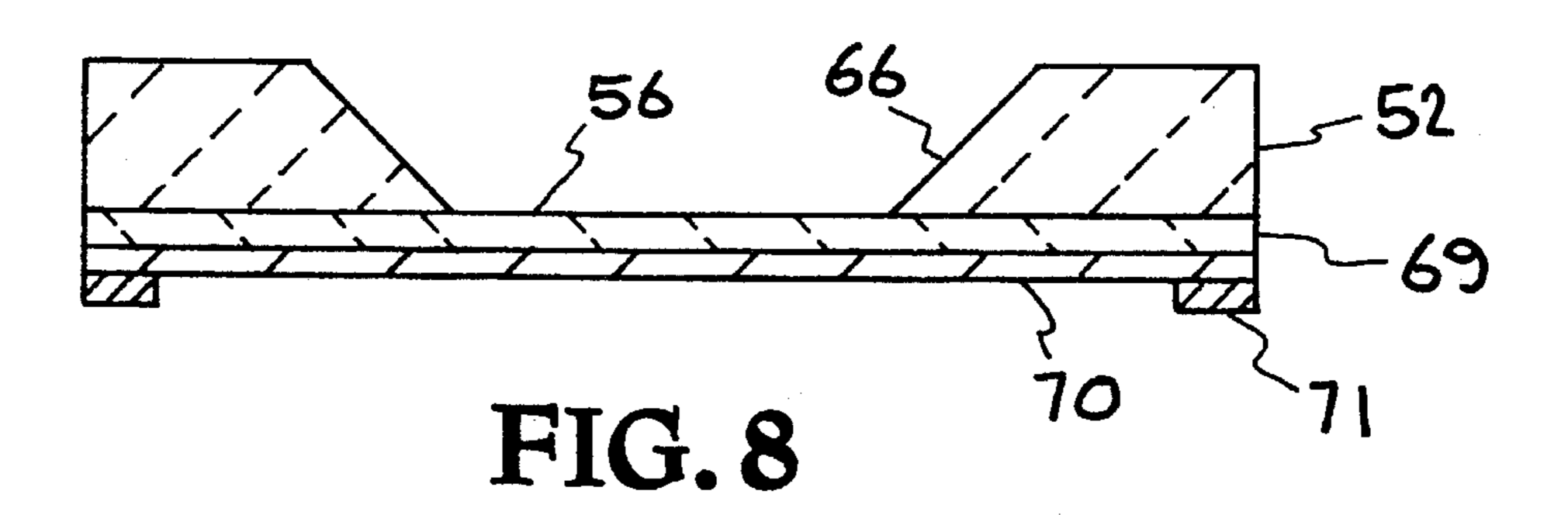
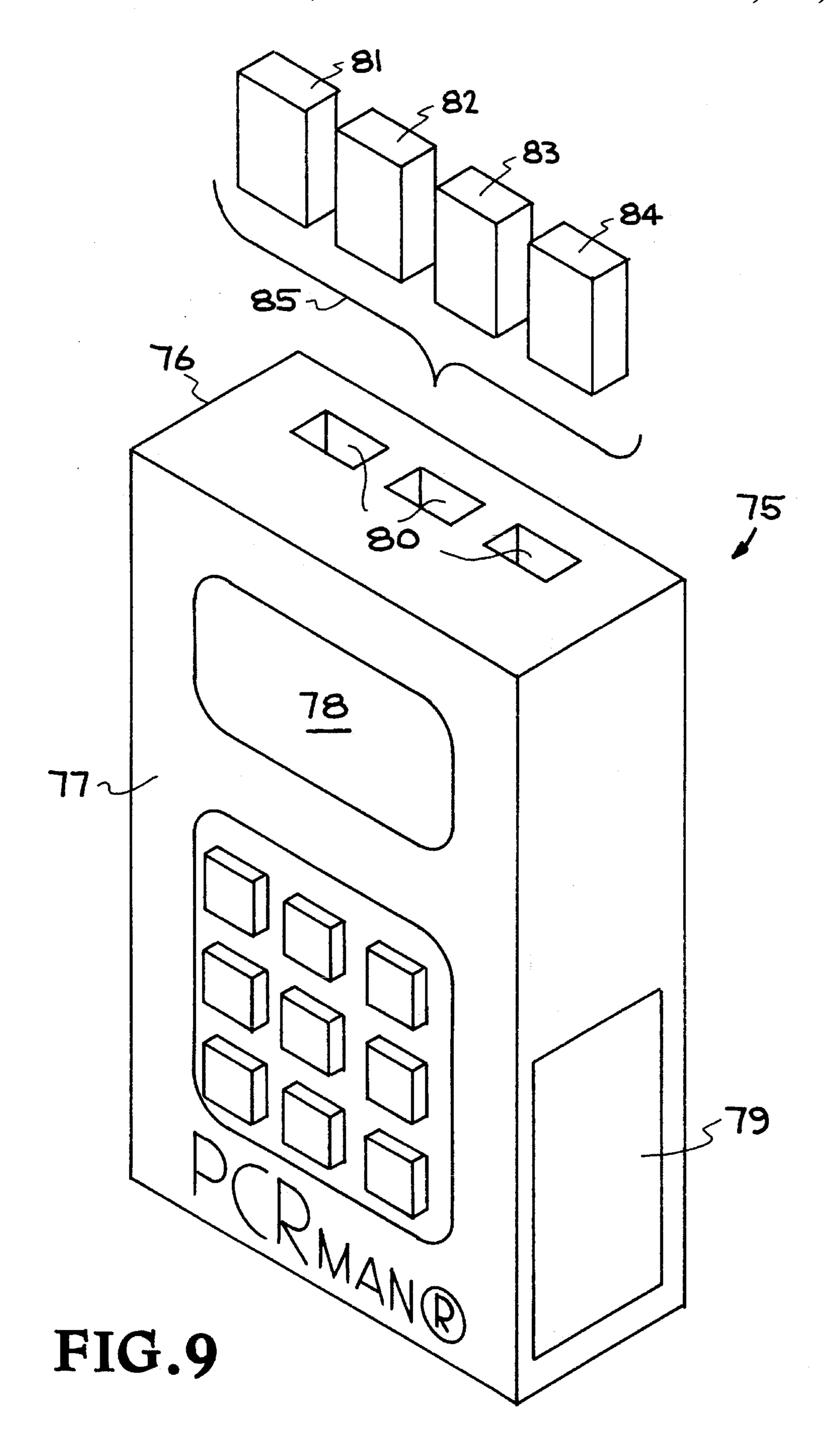


FIG. 7





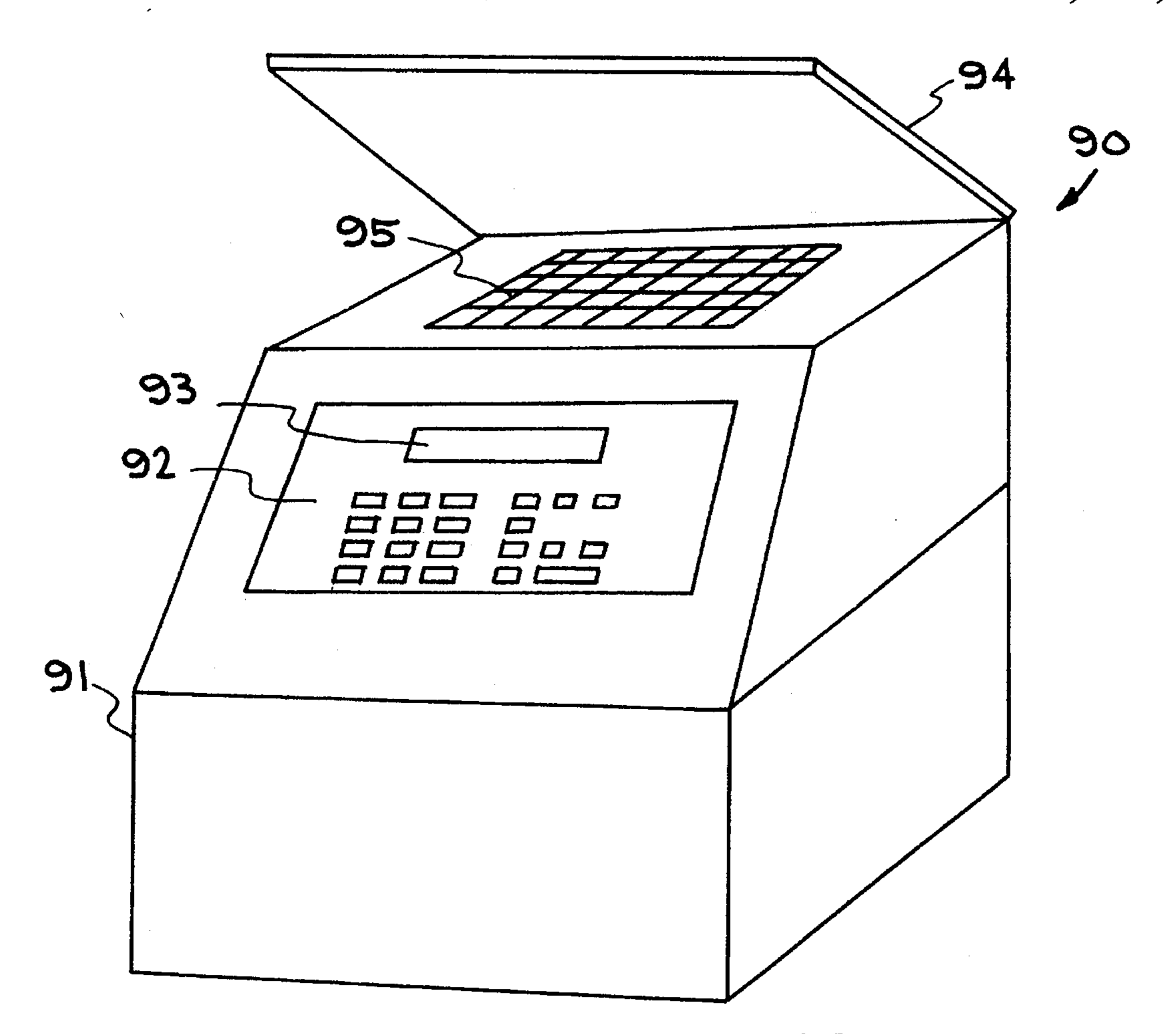
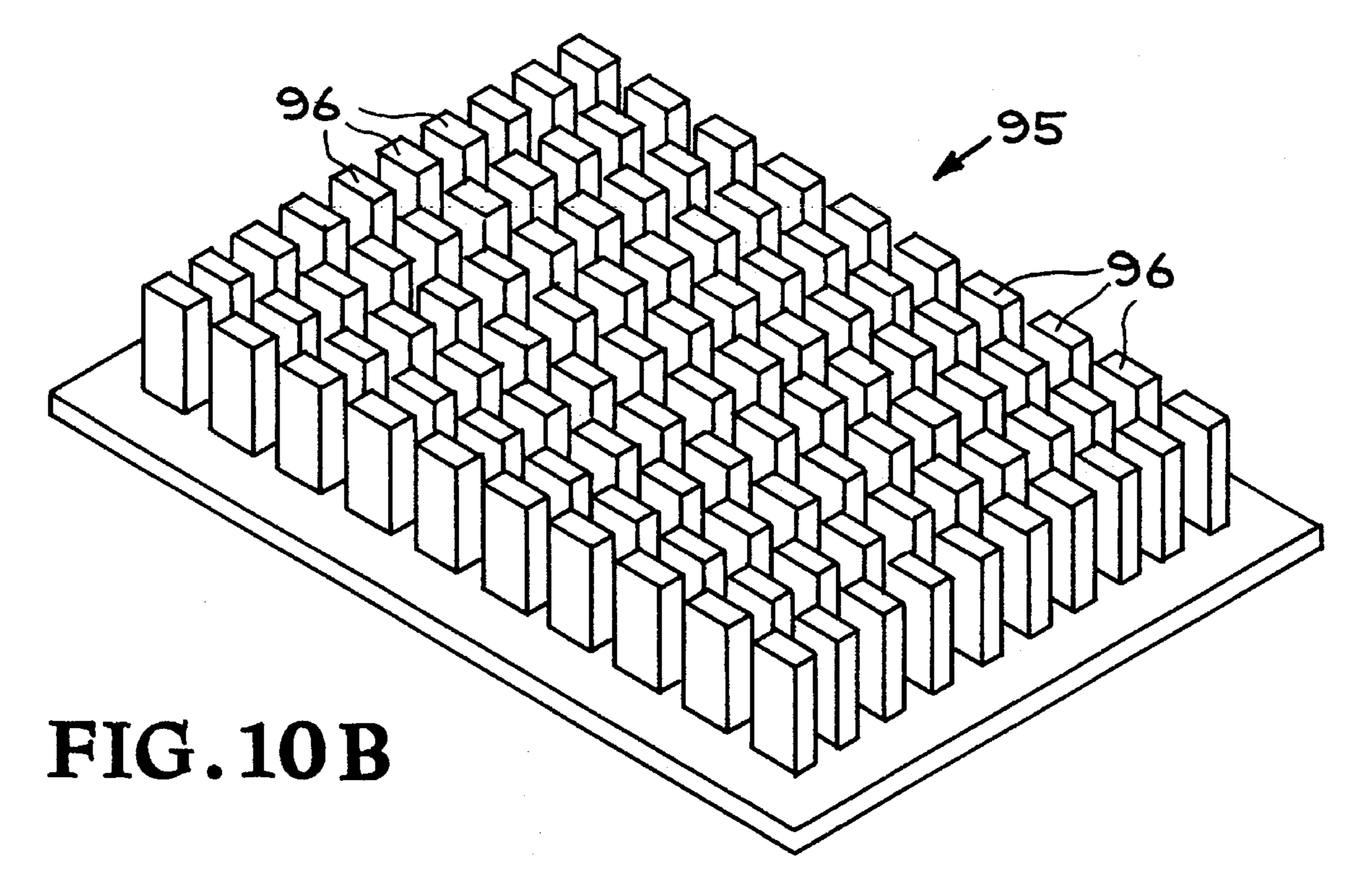
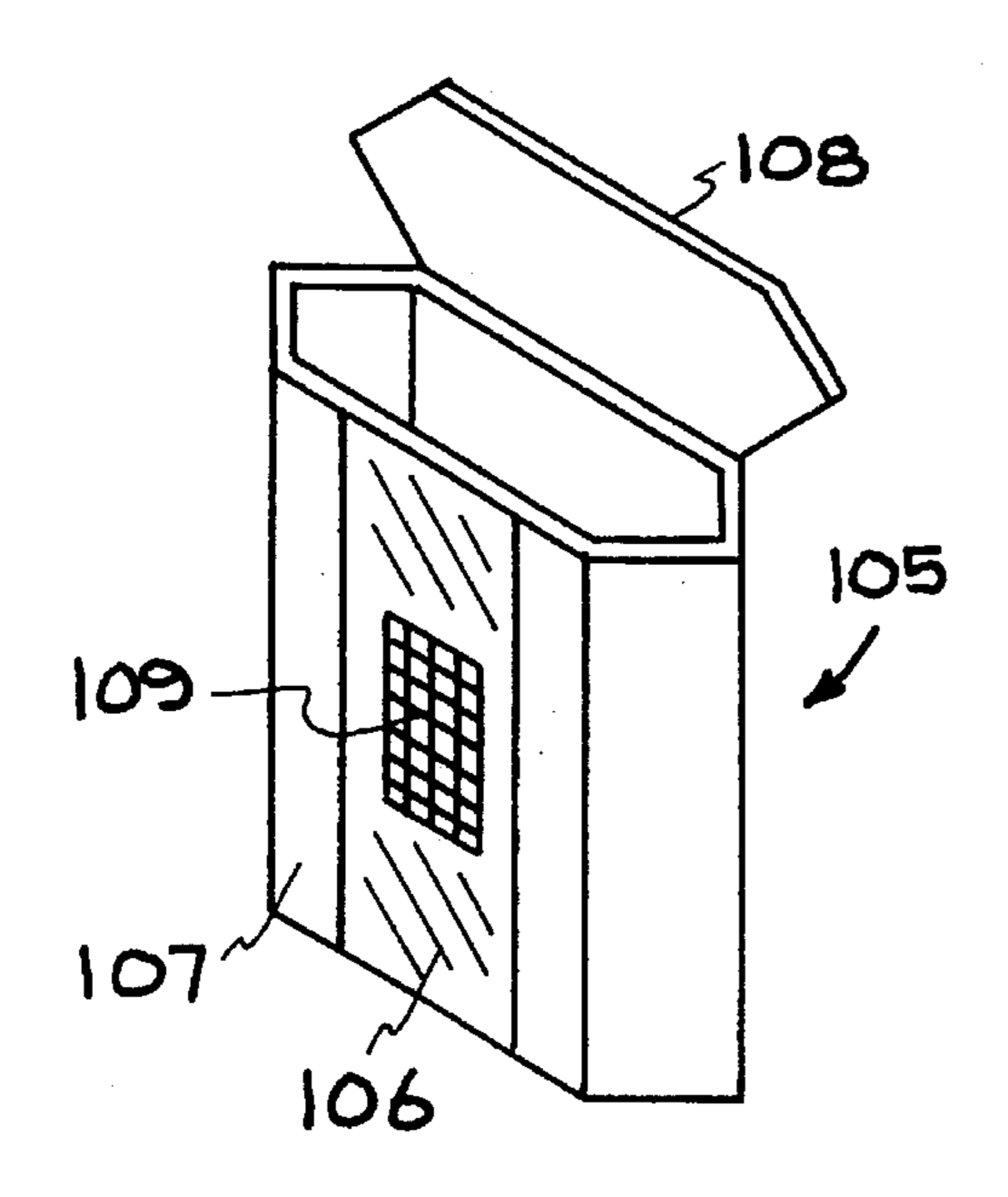


FIG. 10A





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FIG. 12

FIG. 13

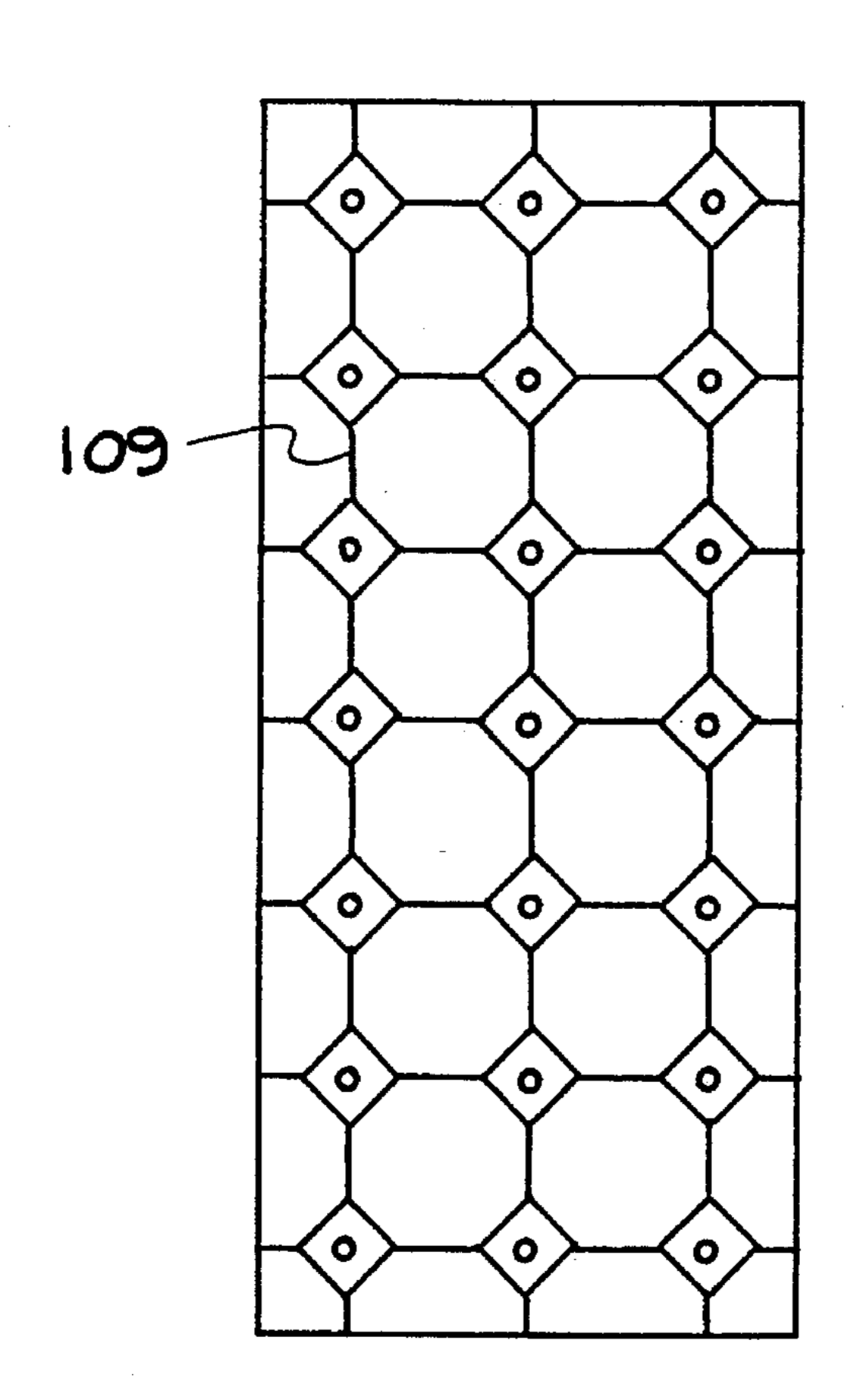
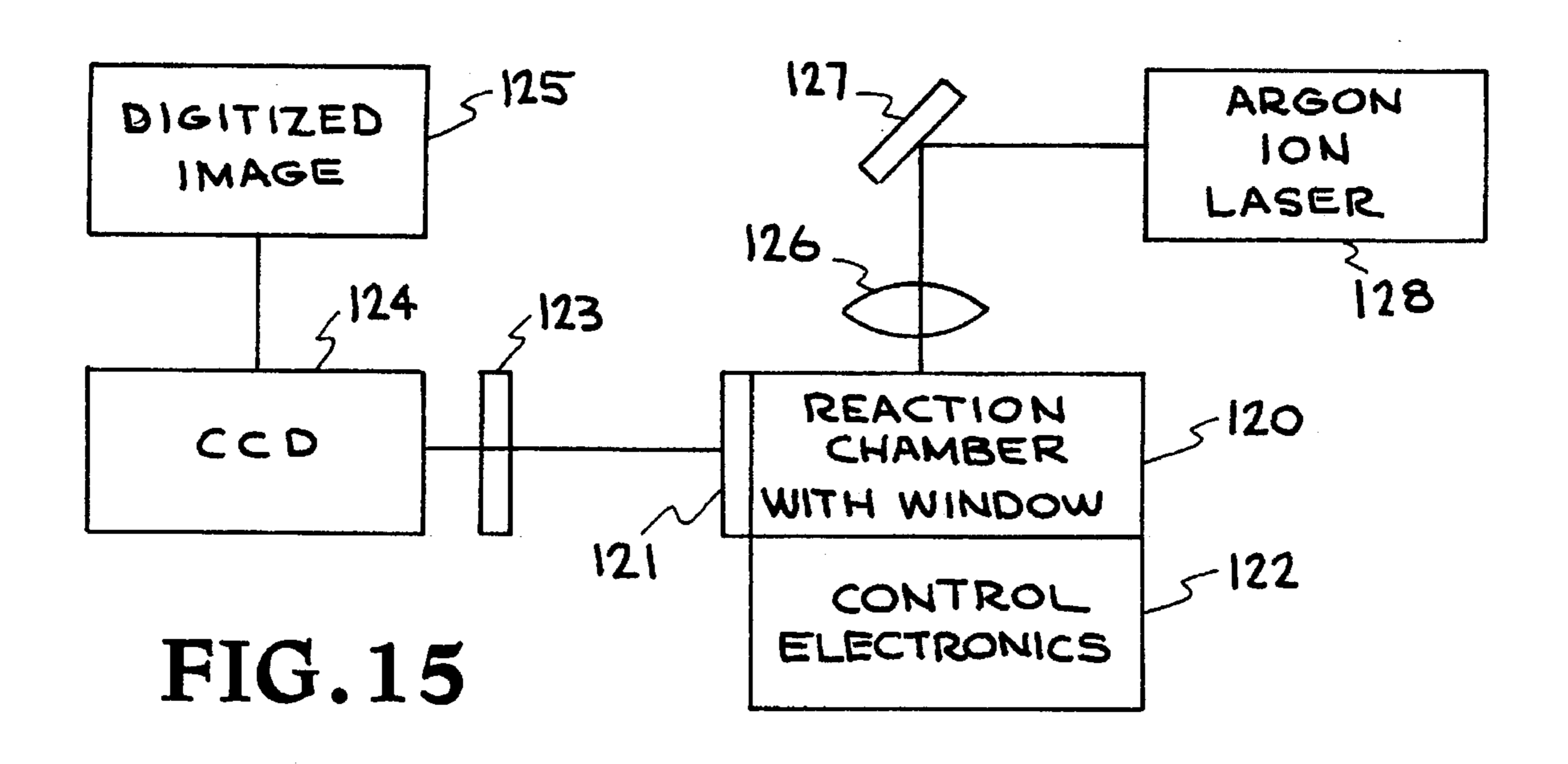
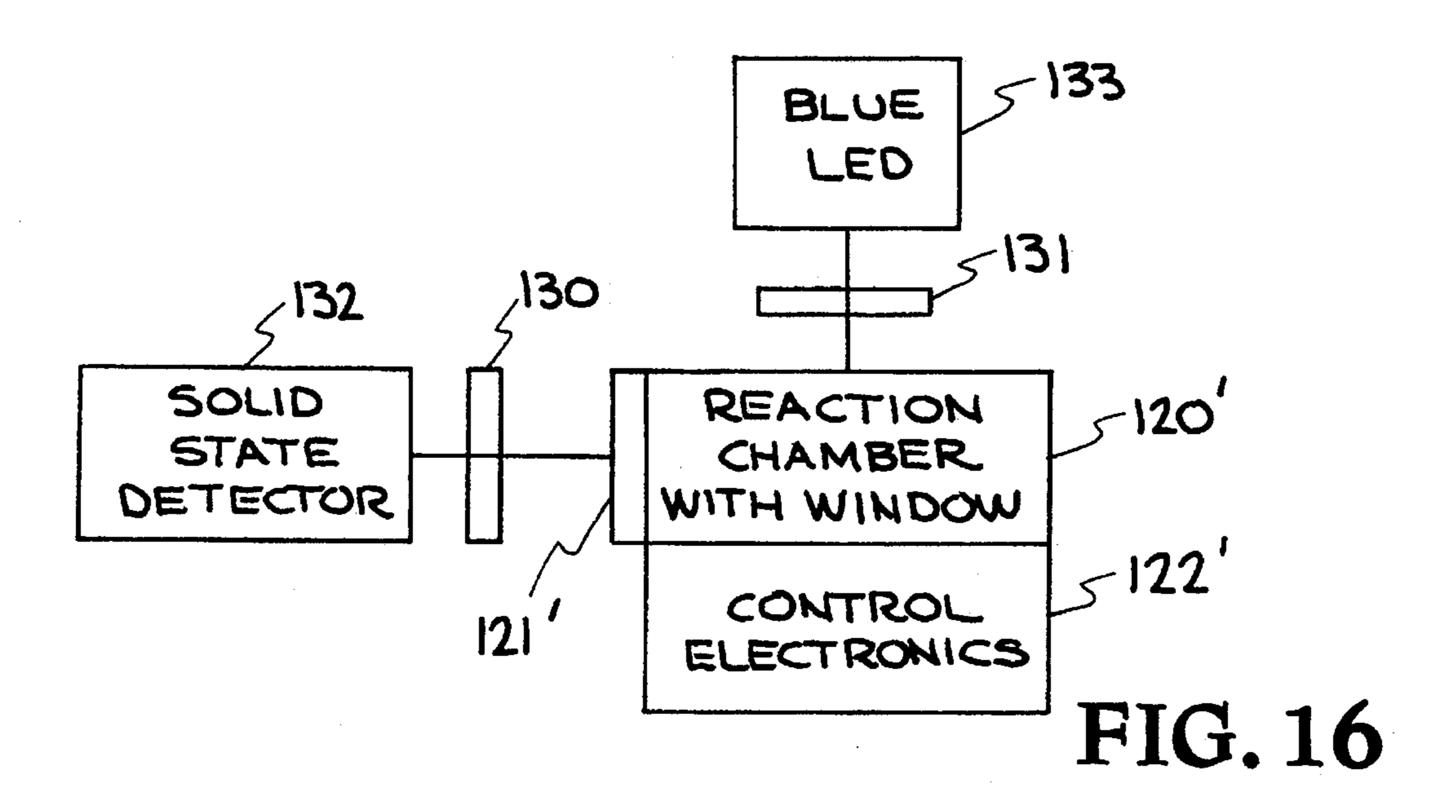
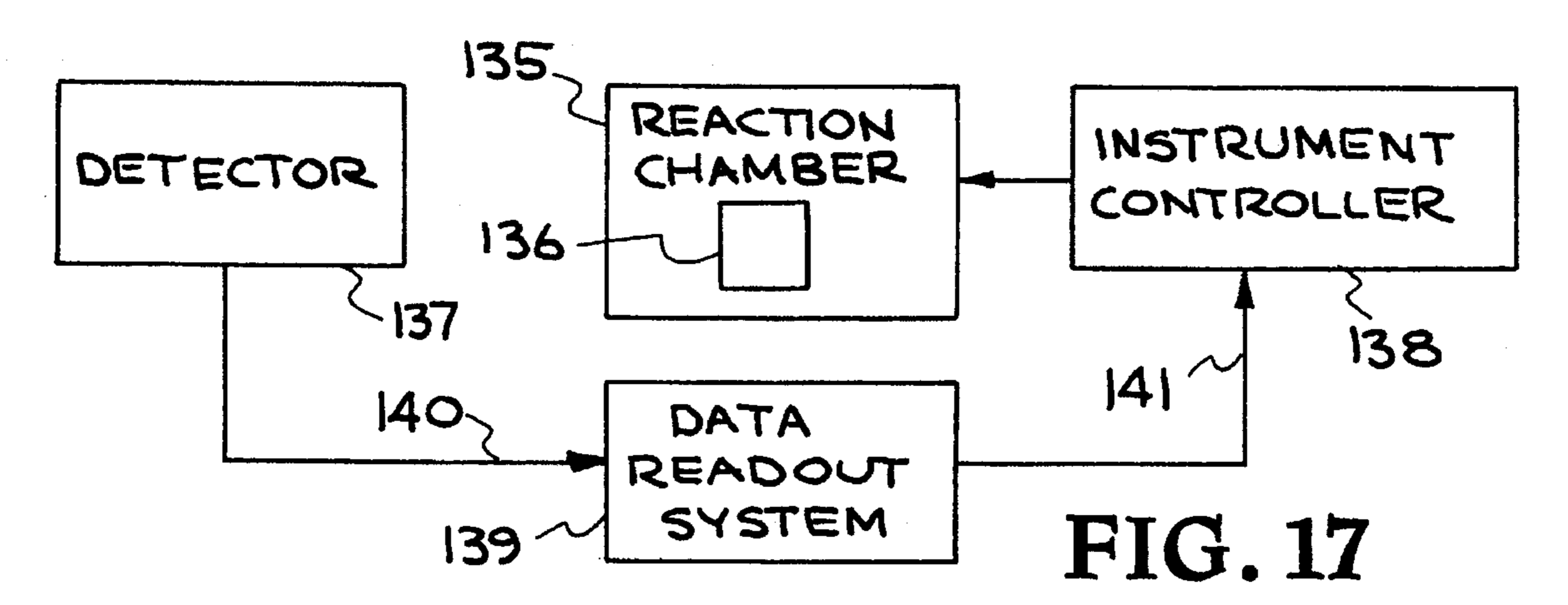


FIG. 14







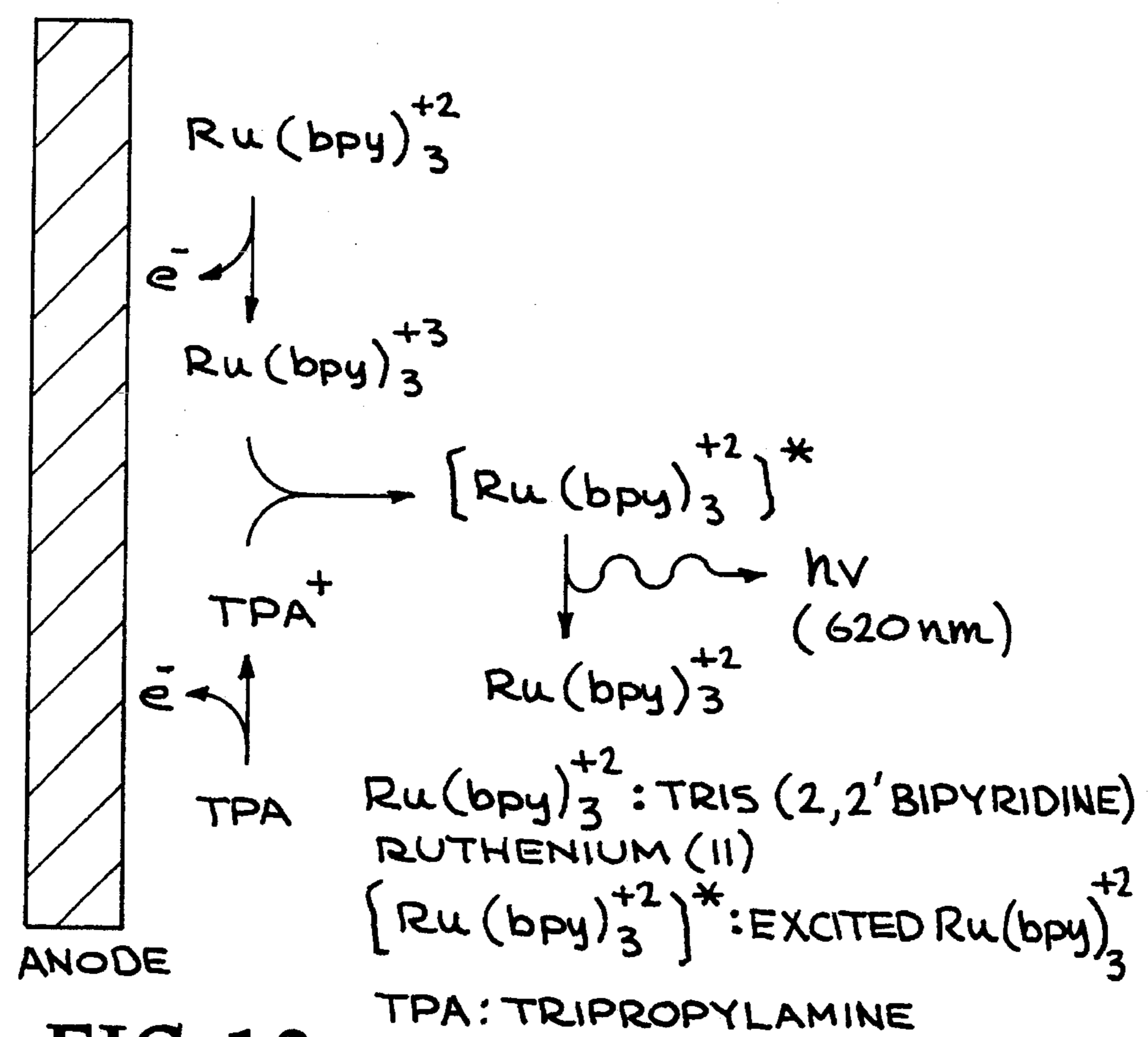
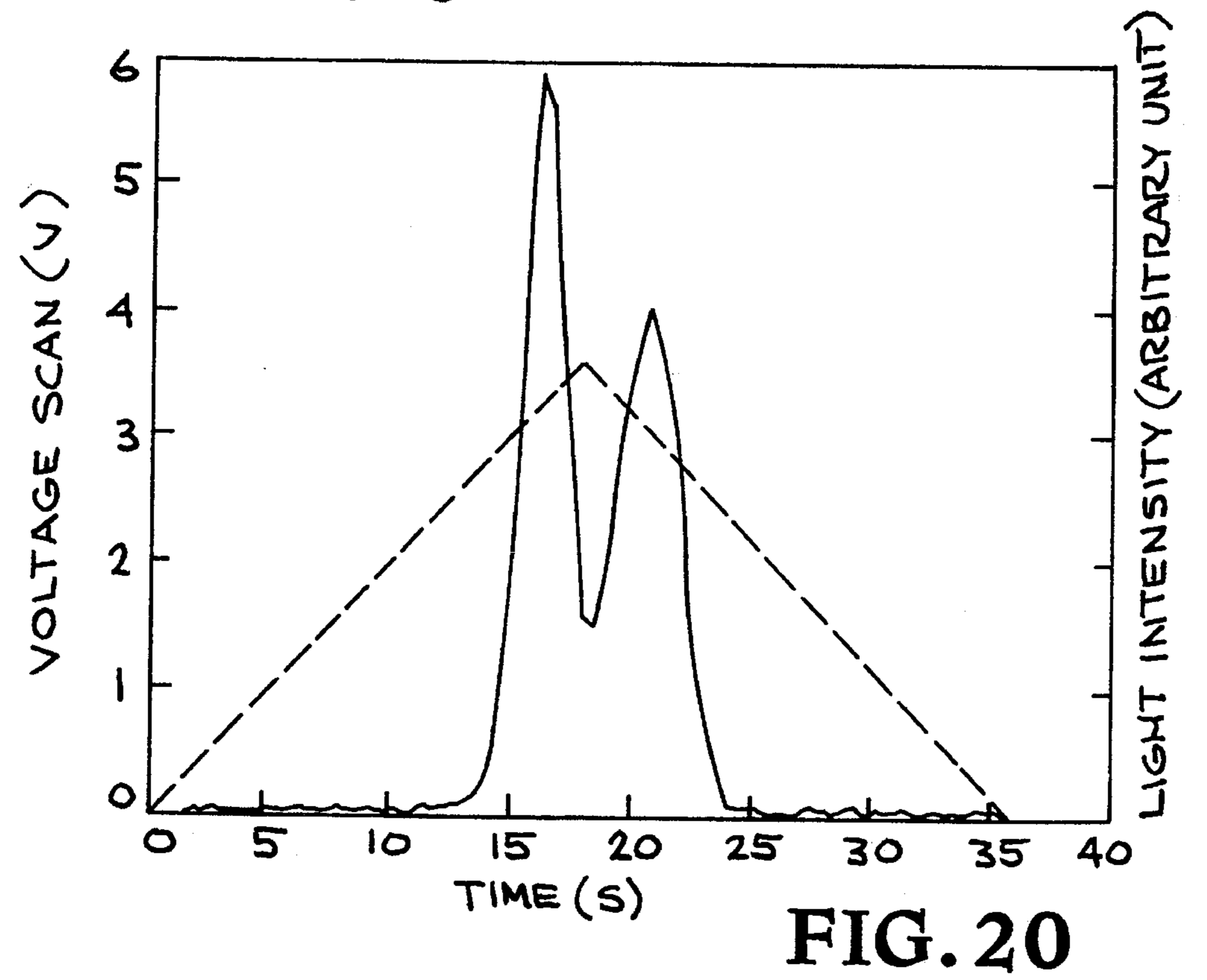


FIG. 18



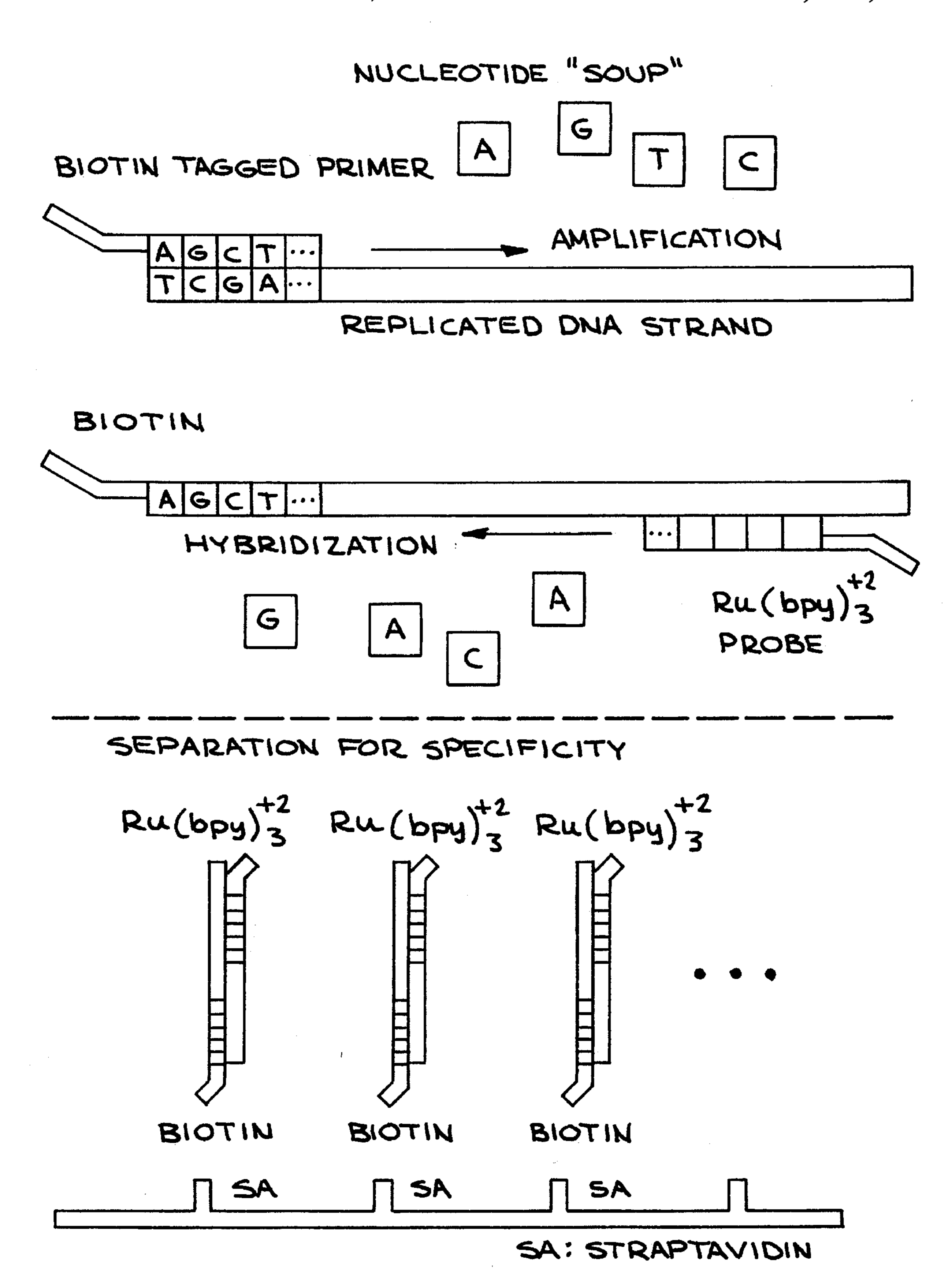
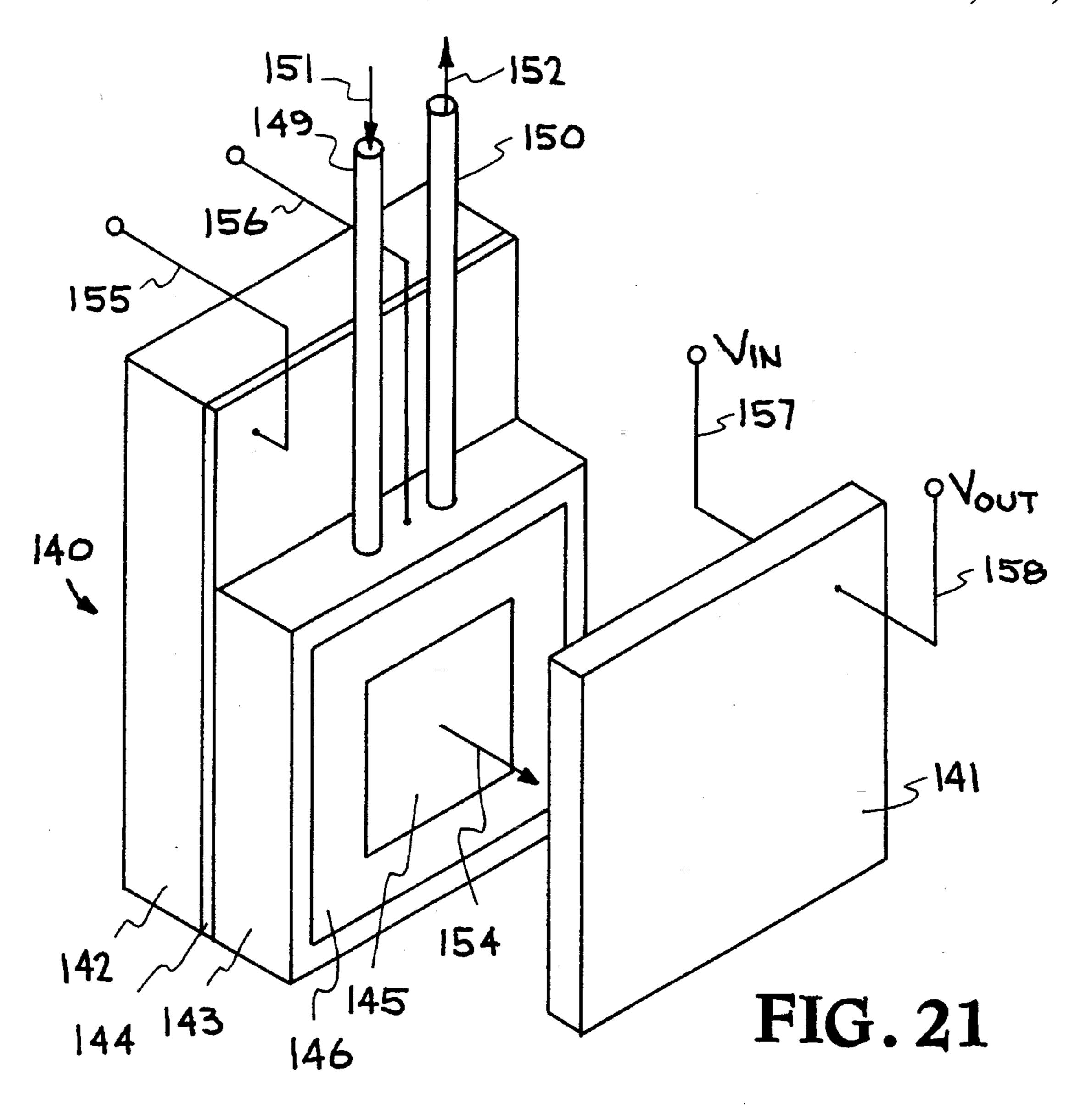
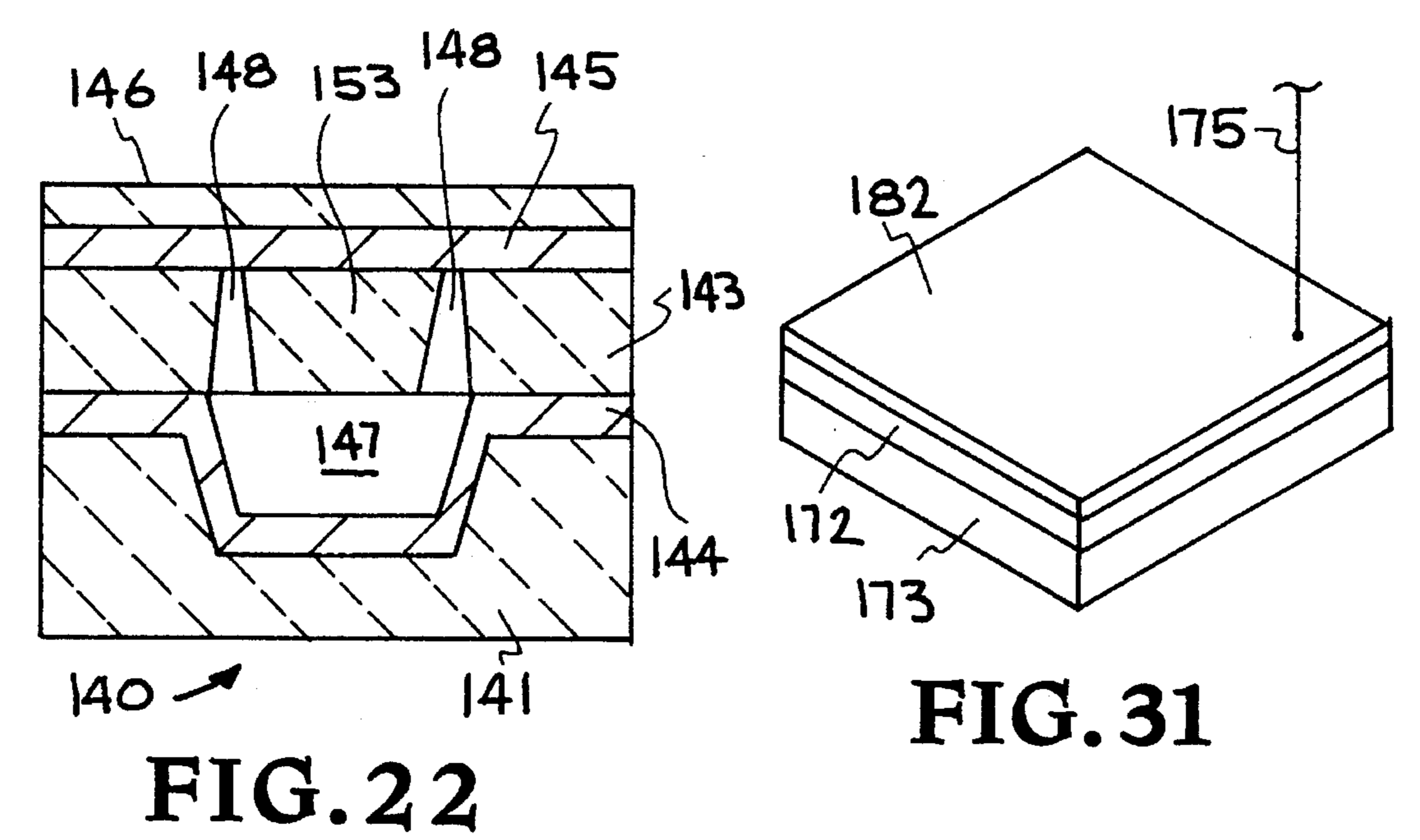


FIG. 19





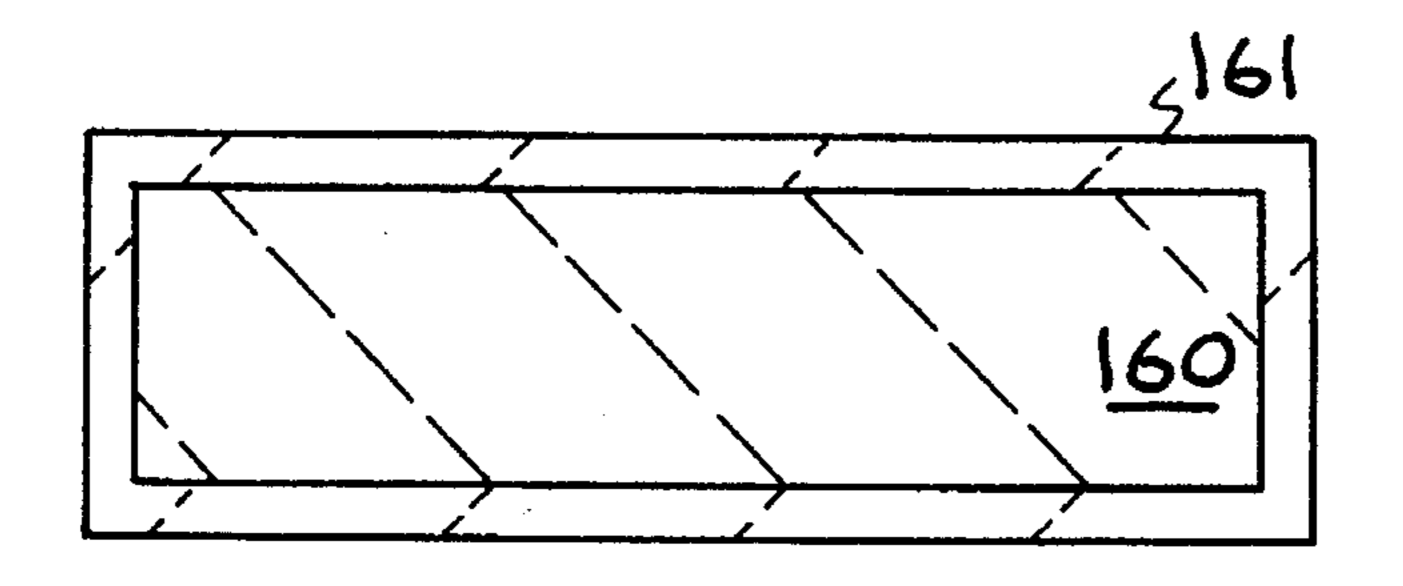
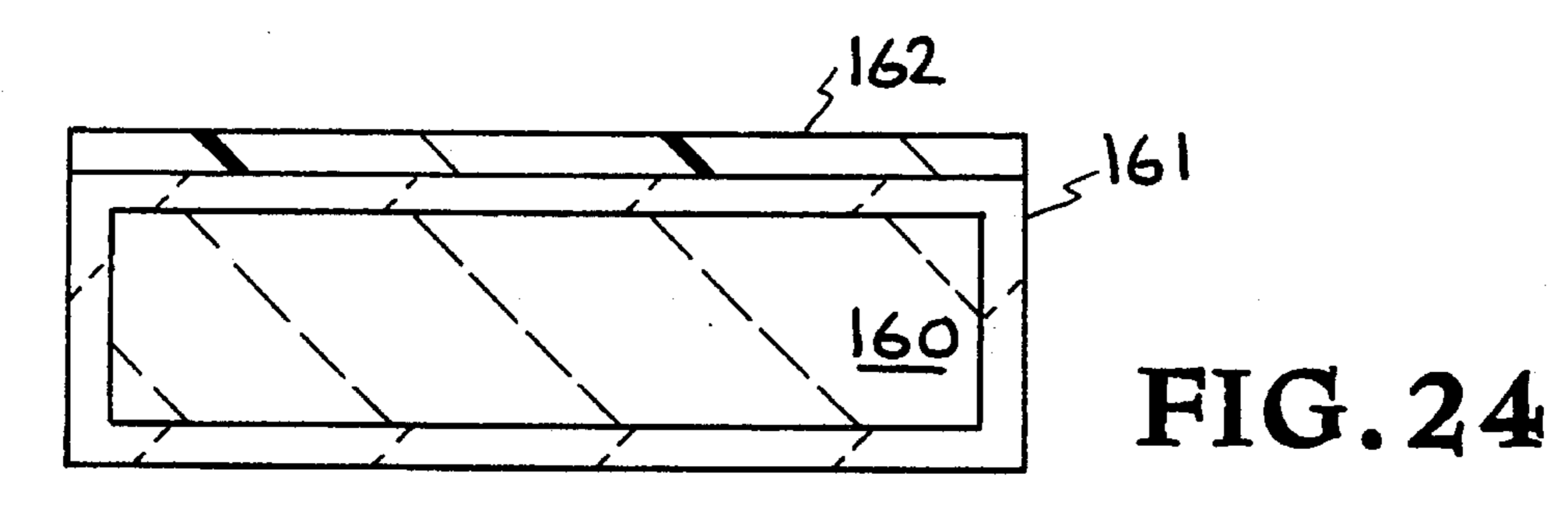


FIG. 23



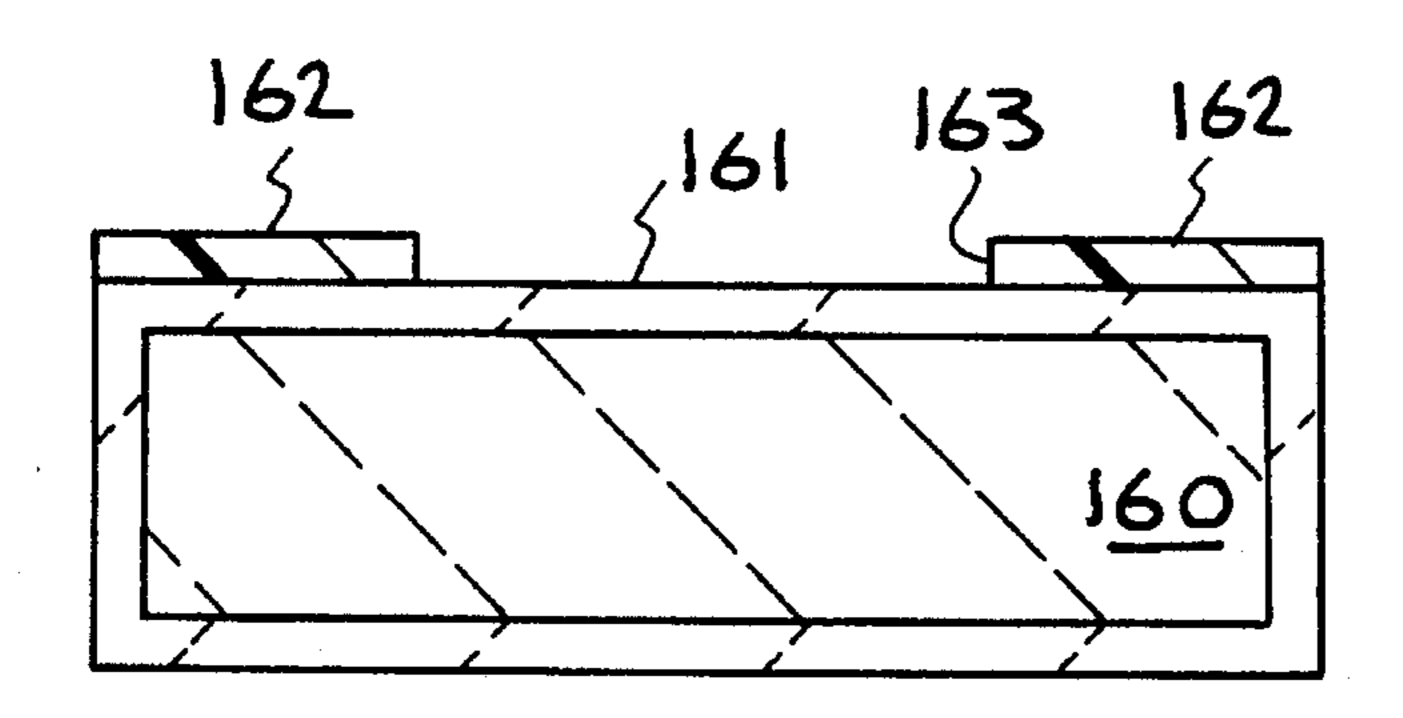


FIG. 25

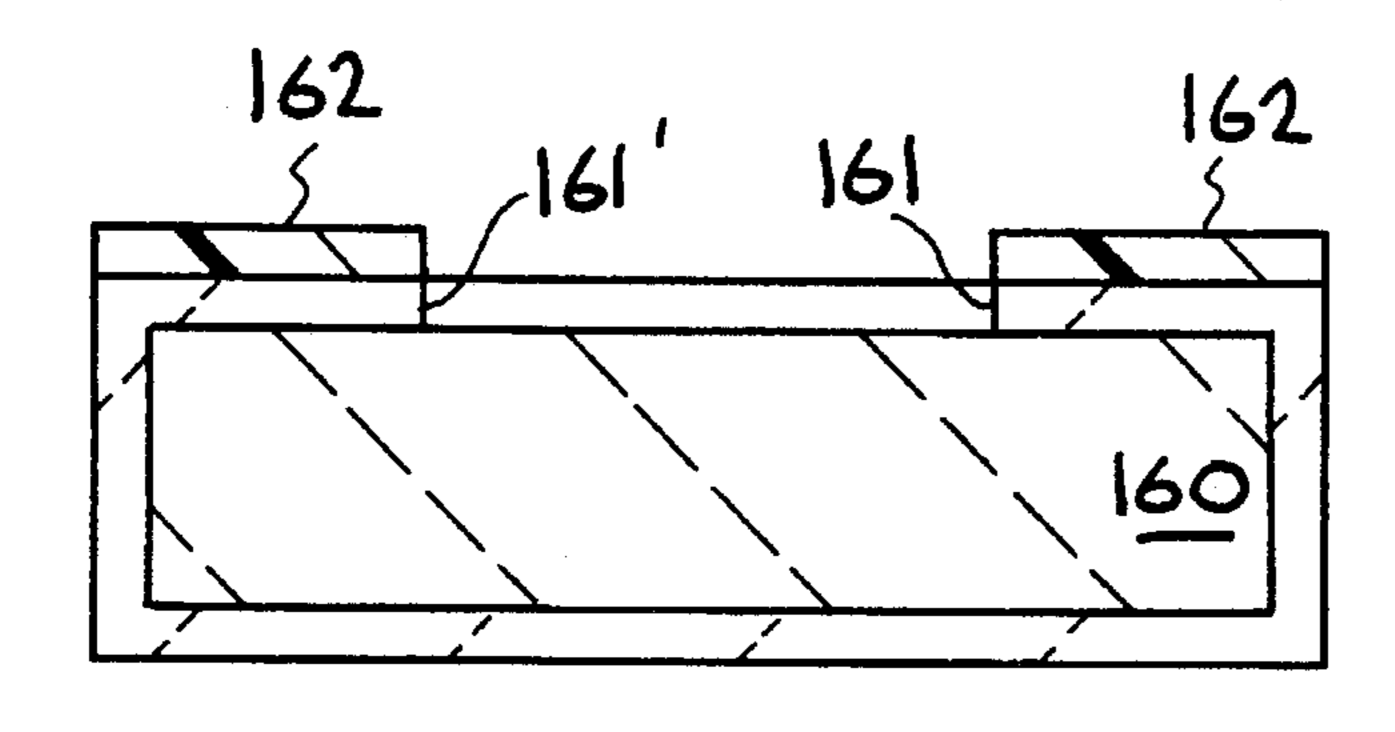


FIG. 26

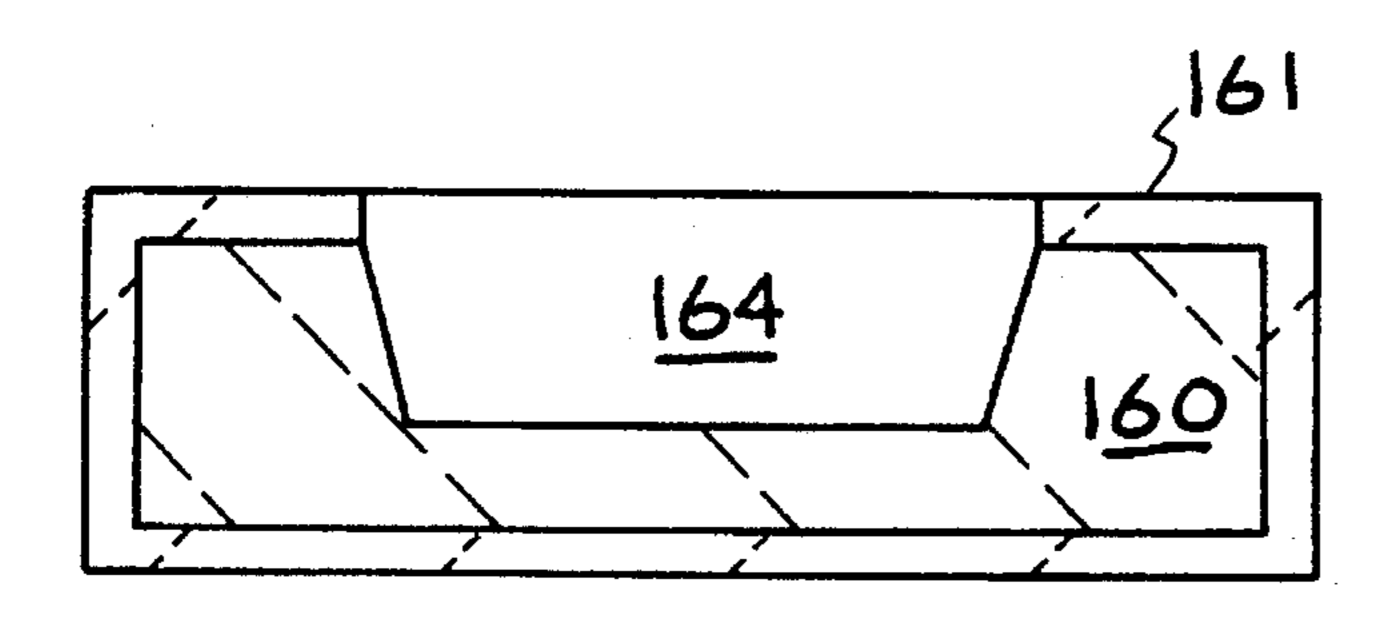
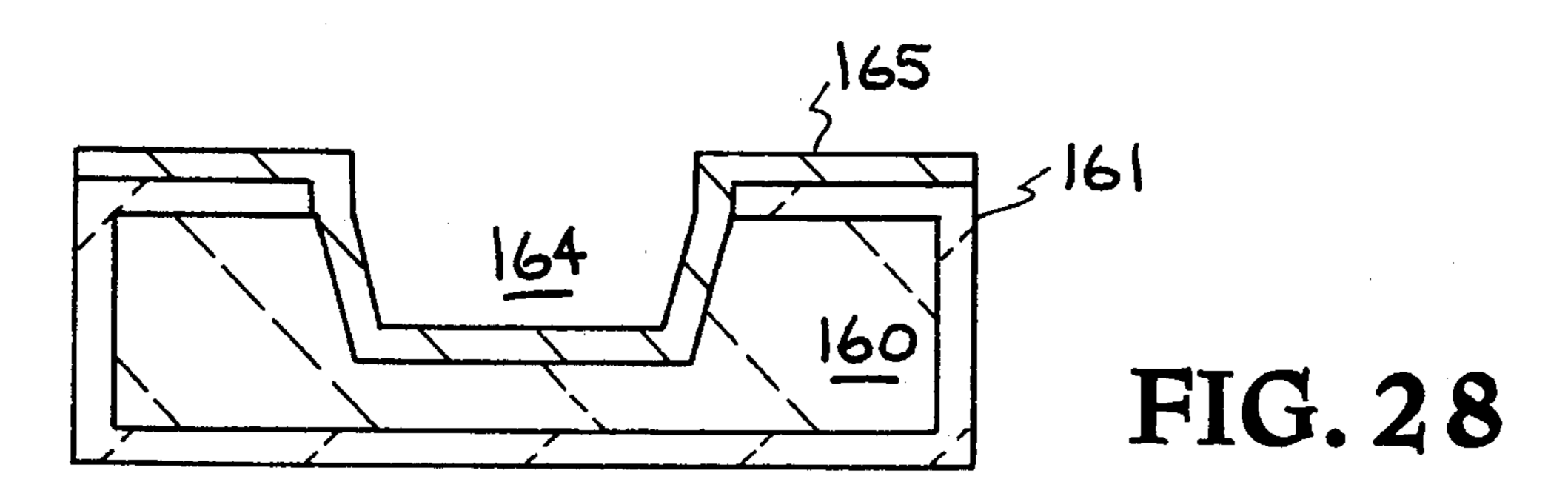
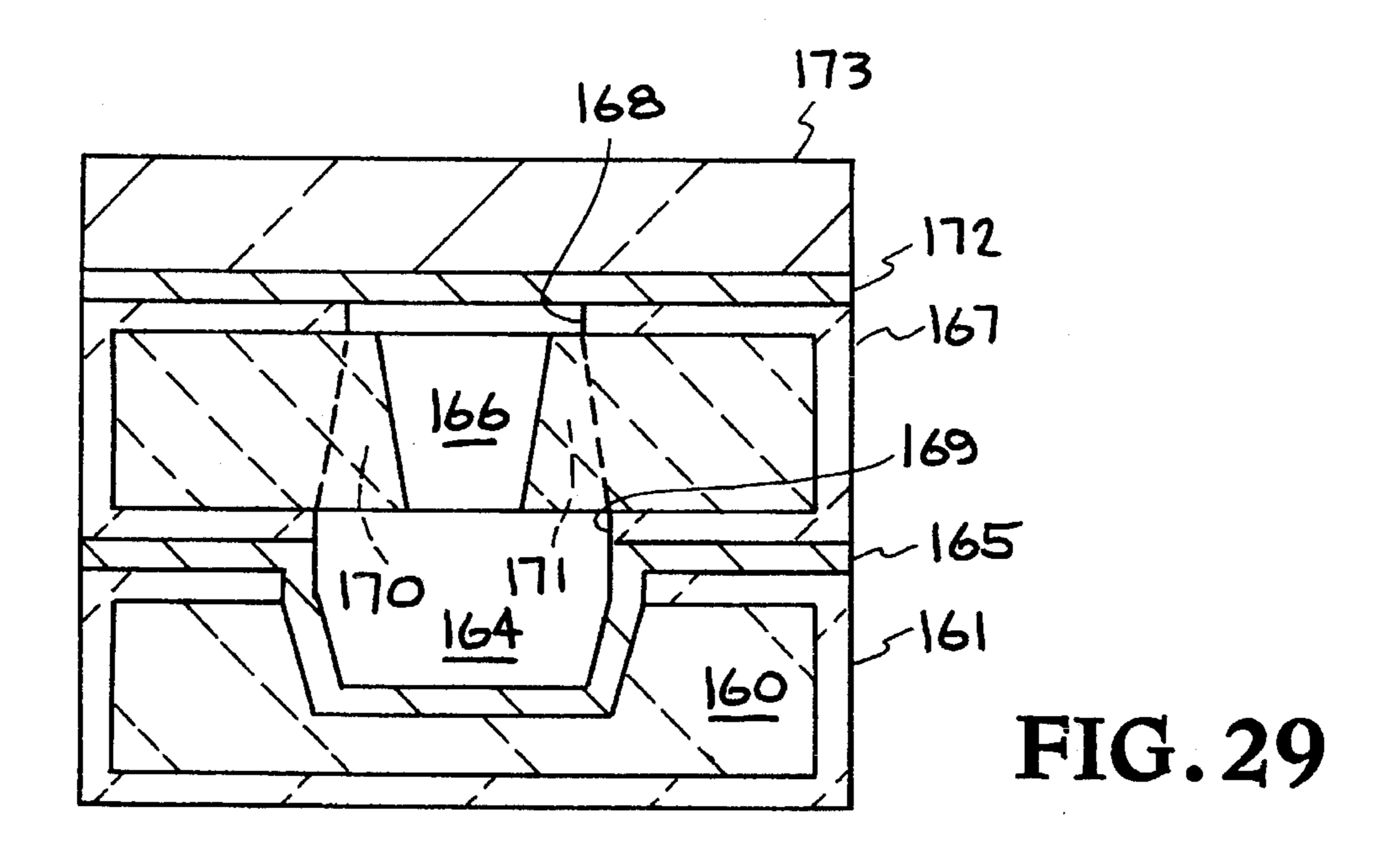
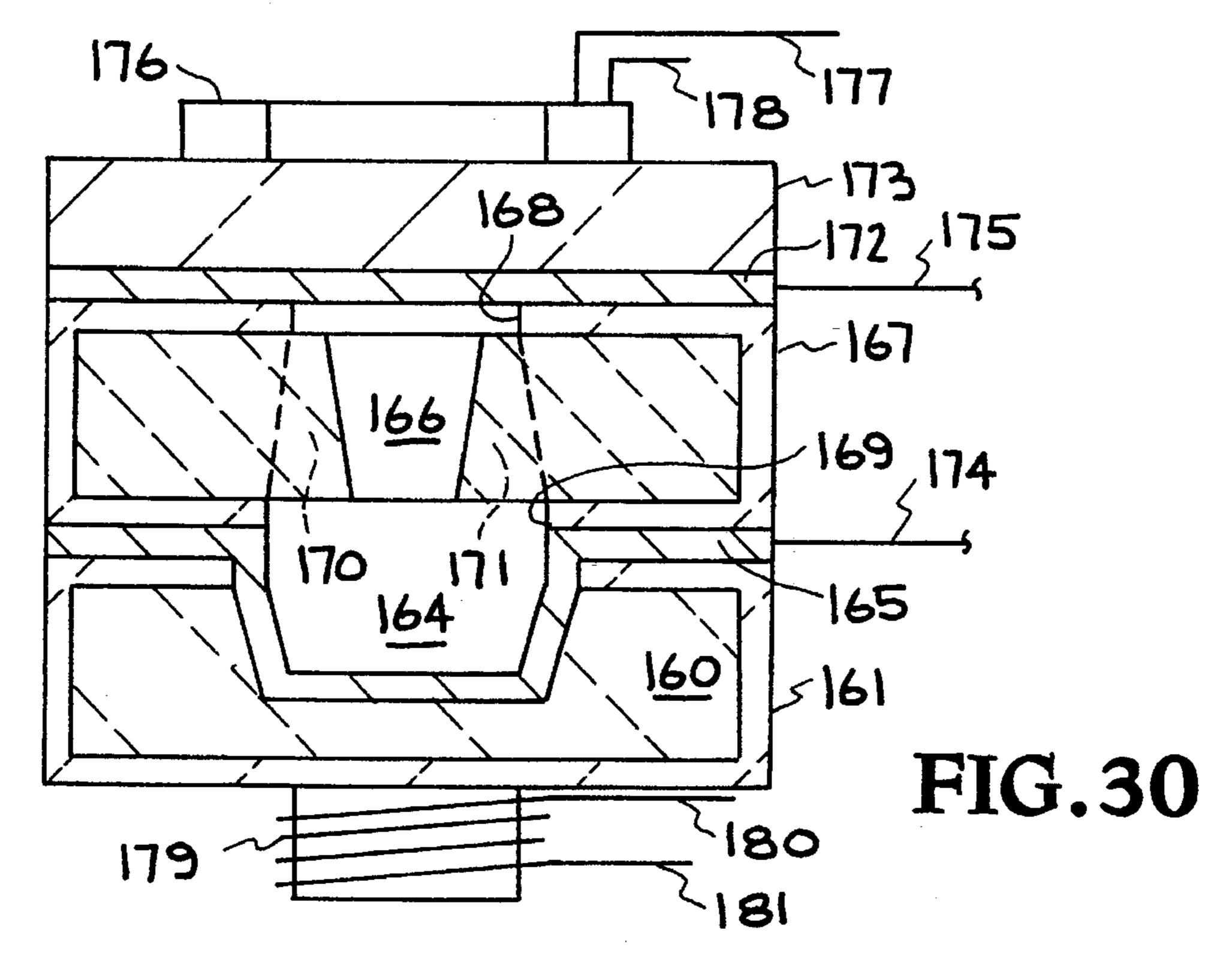


FIG. 27







SILICON-BASED SLEEVE DEVICES FOR CHEMICAL REACTIONS

The United States Government has rights in this invention pursuant to Contract No. W-7405-ENG-48 between the 5 United States Department of Energy and the University of California for the operation of Lawrence Livermore National Laboratory.

BACKGROUND OF THE INVENTION

The present invention relates to instruments for chemical reaction control and detection of participating reactants and resultant products, particularly to integrated microfabricated instruments for performing microscale chemical reactions 15 involving precise control of parameters of the reactions, and more particularly to silicon-based sleeve devices as reaction chambers for chemical reactions and which can be utilized inlarge arrays of individual chambers for a high-throughput microreaction unit.

Current instruments for performing chemical synthesis through thermal control and cycling are generally very large (table-top) and inefficient, and often they work by heating and cooling of a large thermal mass (e.g., an aluminum block). In recent years efforts have been directed to miniaturization of these instruments by designing and constructing reaction chambers out of silicon and silicon-based materials (e.g., silicon, nitride, polycrystalline silicon) that have integrated heaters and cooling via convection through the silicon.

Microfabrication technologies are now well known and include sputtering, electrodeposition, low-pressure vapor deposition, photolithography, and etching. Microfabricated devices are usually formed on crystalline substrates, such as silicon and gallium arsenide, but may be formed on non-crystalline materials, such as glass or certain polymers. The shapes of crystalline devices can be precisely controlled since etched surfaces are generally crystal planes, and crystalline materials may be bonded by processes such as fusion at elevated temperatures, anodic bonding, or field-assisted methods.

Monolithic microfabrication technology now enables the production of electrical, mechanical, electromechanical, optical, chemical and thermal devices, including pumps, valves, heaters, mixers, and detectors for microliter to nanoliter quantities of gases, liquids, and solids. Also, optical waveguide probes and ultrasonic flexural-wave sensors can now be produced on a microscale. The integration of these microfabricated devices into a single systems allows for the batch production of microscale reactor-based analytical instruments. Such integrated microinstruments may be applied to biochemical, inorganic, or organic chemical reactions to perform biomedical and environmental diagnostics, as well as biotechnological processing and detection.

The operation of such integrated microinstruments is easily automated, and since the analysis can be performed in situ, contamination is very low. Because of the inherently small sizes of such devices, the heating and cooling can be extremely rapid. These devices have very low power 60 requirement and can be powered by batteries or by electromagnetic, capacitive, inductive or optical coupling.

The small volumes and high surface-area to volume ratios of microfabricated reaction instruments provide a high level of control of the parameters of a reaction. Heaters may 65 produce temperature cycling or ramping; while sonochemical and sonophysical changes in conformational structures

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may be produced by ultrasound transducers; and polymerizations may be generated by incident optical radiation.

Synthesis reactions, and especially synthesis chain reactions such as the polymerase chain reaction (PCR), are particularly well-suited for microfabrication reaction instruments. PCR can selectively amplify a single molecule of DNA (or RNA) of an organism by a factor of 10⁶ to 10⁹. This well-established procedure requires the repetition of heating (denaturing) and cooling (annealing) cycles in the presence of an original DNA target molecule, specific DNA primers, deoxynucleotide triphosphates, and DNA polymerase enzymes and cofactors. Each cycle produces a doubling of the target DNA sequence, leading to an exponential accumulation of the target sequence.

The PCR procedure involves: 1) processing of the sample to release target DNA molecules into a crude extract; 2) addition of an aqueous solution containing enzymes, buffers deoxyribonucleotide triphosphates (dNTPS), and aligonucleotide primers; 3) thermal cycling of the reaction mixture between two or three temperatures (e.g., 90°–96°, 72°, and 37°–55° C.); and 4) detection of amplified DNA. Intermediate steps, such as purification of the reaction products and the incorporation of surface-bending primers, for example, may be incorporated in the PCR procedure.

A problem with standard PCR laboratory techniques is that the PCR reactions may be contaminated or inhibited by the introduction of a single contaminant molecule of extraneous DNA, such as those from previous experiments, or other contaminants, during transfers of reagents from one vessel to another. Also, PCR reaction volumes used in standard laboratory techniques are typically on the order of 50 microliters. A thermal cycle typically consists of four stages: heating a sample to a first temperature, maintaining the sample at the first temperature, cooling the sample to a second lower temperature, and maintaining the temperature at that lower temperature. Typically, each of these four stages of a thermal cycle requires about one minute, and thus to complete forty cycles, for example, is about three hours. Thus, due to the large volume typically used in standard laboratory procedures, the time involved, as well as the contamination possibilities during transfers of reagents from one vessel to another, there is clearly a need for microinstruments capable of carrying out the PCR procedure.

Recently, the cycling time for performing the PCR reaction has been reduced by performing the PCR reaction in capillary tubes and using a forced air heater to heat the tubes. Also, an integratged microfabricated reactor has been recently developed for in situ chemical reactions, which is especially advantageous for biochemical reactions which require high-precision thermal cycling, particularly DNAbased manipulations such as PCR, since the small dimensions of microinstrumentation promote rapid cycling times. This microfabricated reactor is described and claimed in copending U.S. application Ser. No. 07/938,106, filed Aug. 31, 1992, entitled "Microfabricated Reactor", assigned to the same assignee. Also, an optically heated and optically interrograted micro-reaction chamber, which can be utilized, for example, in the integrated microfabricated reactor of the above-referenced copending application Ser. No. 07/938, 106, has been developed for use in chemical reactors, and is described and claimed in copending U.S. application Ser. No. 08/489,918, filed Jun. 13, 1995, entitled Diode Laser Heated Micro-Reaction Chamber With Sample Detection Means", assigned to the same assignee.

The present invention is directed to a particular geometry of silicon-based micro-reactors that have shown to be very

efficient in terms of power and temperature uniformity. The micro-reactor of this invention, which is broadly considered as a silicon-based sleeve device for chemical reactions, can be effectively utilized in either of the reactor systems of the above-referenced copending applications. The present 5 invention utilizes doped polysilicon for heating and bulk silicon for convective cooling. The present invention allows the multi-parameter, simultaneous changing of detection window size, in situ detection, reaction volumes, thermal uniformity, and heating and cooling rates. In addition, it 10 enables the use of large arrays of the individual reaction chambers for a high-throughput microreaction unit.

SUMMARY OF THE INVENTION

It is an object of the present invention to provide an improved chemical reaction chamber.

A further object of the invention is to provide a siliconbased sleeve device for chemical reactors.

A further object of the invention is to provide a chemical 20 reaction chamber that combines to use of doped polysilicon and bulk silicon.

A further object of the invention is to provide chemical reaction chambers that combines the use of doped polysilicon and bulk silicon to provide flexibility in thermal and optical properties allowing the implementation into small and large instruments.

Another object of the invention is to provide a silicon-based reaction sleeve that combines a criticial ratio of silicon and silicon nitride to the volume of material to be heated (e.g., liquid) in order to provide uniform heating, yet low power requirement.

Another object of the invention is to provide a siliconbased reaction sleeve that will allow the introduction of a secondary tube (e.g., plastic) into the reaction sleeve that contains the reaction mixture, thereby eleviating any potential materials incompatiblity issues.

Another object of the invention is to provide an array of individual reaction chambers for a high-throughput microre- 40 action unit.

Another object of the invention is to provide a hand-held instrument that uses silicon-based sleeve-type reaction chambers with integrated heaters.

Another object of the invention is to provide a reaction chamber with automated detection and feedback control.

Another object of the invention is to provide for artificial intelligence control of reactions in a reaction chamber.

Another object of the invention is to provide pulse-width 50 modulation as a feedback control for reaction chamber.

Other objects and advantages of the present invention will become apparent from the following description and the accompanying drawings. Basically, the invention is a silicon-based sleeve for chemical reactions. The invention 55 encompasses a chemical reaction chamber that combines the use of polysilicon for heating and bulk silicon for convective cooling. The reaction sleeve combines a critical ratio of silicon and silicon nitride to the volume of material to be heated in order to provide uniform heating, yet low power 60 requirements. The reaction sleeve also allows for the introduction therein of a secondary tube that contains the reaction mixture thereby eleviating any potential materially incompatibility issues. The present invention is an extension of the above-referenced integrated micofabricated reactor of 65 above-referenced copending application Ser. No. 07/938, 106 and the above-references optically integrated micro4

reaction chamber of above-referenced copending application Ser. No. 08/489,819. The silicon-based sleeve reaction chamber can be utilized in chemical reaction systems for synthesis or processing of organic, inorganic, or biochemical reactions, such as the polymerase chain reaction (PCR) and/or other DNA reactions (such as the ligose chain reaction), or other synthetic, thermal-cycling-based reactions.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 shows a partial cut-away perspective view of a microfabricated chemical reaction instrument mounted in a power source / control apparatus.

FIG. 2 is a schematic of the reaction instrument of FIG. 1

FIG. 3 schematically illustrates a heating and detection arrangement for a microfabricated reaction chamber.

FIG. 4 illustrates an embodiment of a microfabricated silicon-based sleeve reaction chamber made in accordance with the present invention.

FIG. 5 is an array of the sleeve reaction chambers of FIG. 4 operatively connected to a microelectrophoresis array.

FIG. 6 is an enlarged end view of another embodiment of a sleeve microreaction chamber similar to FIG. 4.

FIG. 7 illustrates in cross-section embodiment of an enlarged section of FIG. 6 using an isolated heater version, fixed window.

FIG. 8 illustrates in cross-section another embodiment of the same enlarged section of FIG. 6 using a non-isolated heater version variable window.

FIG. 9 is a view of a hand-held instrument (PCR man) which utilizes the reaction chambers of FIG. 6 as inserts to change reactions.

FIGS. 10A and 10B illustrate a thermal cycling instrument utilizing several hundreds of individually-controlled silicon-based microreaction chambers.

FIG. 11 illustrates a schematic representation of high-throughput DNA amplification, sample-handling, and electrophoreseis system.

FIG. 12 is an embodiment of an insert/lining for a reaction chamber with optical window, with the top/cover open.

FIG. 13 illustrates external filling of a reaction chamber insert/liner.

FIG. 14 illustrates immobilized reagents/probes for detection of specific products directly on windows or within reaction fluid as "test strip" detected optically in the hand held instrument (PCR man) of FIG. 9.

FIGS. 15 and 16 schematically illustrate optical detection systems for use with the microreaction chambers of FIG. 6.

FIG. 17 schematically illustrates the use of integrated detection for an artificial intelligent feedback system.

FIG. 18 is a diagram showing the electrochemical oxidation and chemical reduction reactions for tris (2,2'bipyridyl) ruthenium (II) (TBR) and tripropylamine (TPA).

FIG. 19 illustrates a method for tagging and separating DNA for detection and quantification by electrochemiluminescence (ECL).

FIG. 20 illustrates cell voltage and ECL intensity versus time, with the voltage being increased, then decreased.

FIG. 21 illustrates an embodiment of a micromachined ECL cell with a thin film anode, and an associated photodiode detector.

FIG. 22 is an enlarged cross-sectional view of the ECL cell of FIG. 21 with the electrical leads.

FIGS. 23–30 illustrate the fabrication process for producing an ECL cell, as illustrated in FIG. 21.

FIG. 31 illustrates an embodiment using Al on ITO on glass which reduces resistance of the ITO electrode.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is a micro-fabricated silicon-based sleeve chemical reaction chamber that combines heaters, such as doped polysilicon for heating and bulk silicon for conventive cooling. The microreaction chambers can be used in an array for a high-throughput microreaction unit, or in a hand-held unit. It combines a critical ratio of silicon and silicon nitride to the volume of material to be heated (e.g., liquid) in order to provide uniform heating, yet low power requirements. It also will allow the introduction of a secondary tube (e.g., plastic) into the reaction sleeve that contains the reaction mixture thereby alleviating any potential materials incompatibility issues. The present invention utilizes a particular geometry of silicon-based micro-reactors that have been shown to be very efficient in terms of power and temperature uniformity. The particular embodiment of the microfabricated reactor described has been experimentally used as a thermal cycling instrument for use in the polymerase chain reaction (PCR) and other chemical reactions, and has shown to be superior to present commercial instruments on thermally-driven chemical reactors. The silicon-based sleeve reaction chamber of this invention can be utilized in place of the reaction chamber of the microfabricated system of above-referenced copending application Ser. No. 07/938,106; and can be utilized with the integrated heater and detection arrangement of above-referenced copending application Ser. No. 08/489,819; and thus constitutes an extension of the microfabricated chemical reaction systems in these copending applications.

To provide an understanding of a microfabricated chemical reaction instrument and the integrated heating/detection arrangement, prior to the description of the embodiment of the sleeve reaction chamber of the present invention, a description is set forth of a microfabricated chemical reactor and an integrated heating/detection arrangement of the two-referenced copending applications.

FIG. 1 illustrates an embodiment of a microfabricated chemical reaction instrument generally indicated at 10, shown above a recessed section thereof, indicated generally at 11, in a power source/control system of the microfabricated reaction instrument, generally indicated at 12. A hypodermic needle 13 is shown inserting a sample through a silicone rubber window 14 into the reaction instrument 10. The reaction is controlled and powered by: induction coupling, such as that between coil L_{CL} in the instrument 10 and a magnetic coil 15; by capacitive coupling, such as that between the plates of capacitor C_3 and plates 16 and 17; and 150 by electromagnetic coupling between a resonant circuit, see FIG. 152, in instrument 103 and a radio frequency antenna 153.

A schematic of the instrument 10 of FIG. 1 is illustrated in FIG. 2, and comprises three reagent chambers 19, 20 and 21, which, for example, may contain the DNA primers, the 60 polymerase, and the nucleotides and any detection-tag molecules, such as magnetic beads. The target DNA molecule is placed in reagent chamber 19 by insertion of a hypodermic needle 13 (FIG. 1) or the like through a silicone rubber or other type material window 14. The reactants chambers 19, 65 20 and 21 are respectively connected by channels 22, 23, and 24, having narrow midsections, not shown, to a reaction

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chamber 25. Typically the chambers 19–21 and 25 have a volume ranging from microliter to nanoliters. The channels 22–24 are equipped with Lamb-wave pumps LW₁, LW₂ and LW₃, respectively, for pumping reactants in chambers 19–21 through channels 22–24 in the direction of the arrows into reaction chamber 25. The Lamb-wave pumps may be located on any wall, or on multiple walls, of the channels 22–24. The Lamb-wave pumps LW₁, LW₂, and LW₃ are connected respectively to capacitors C₁, C₂, and C₃. The surface tension across the narrow midsections of the channels 22–24 prevents the reactants in chambers 19–21 from flowing into reaction chamber 25 until pumping is initiated. The inner surfaces of the channels 22–24 may be treated to raise the surface tension thereby further inhibiting flow of the reagents when the Lamb-wave pumps are not activated.

The reaction chamber 25 may be equipped with a Lambwave transducer LW_C and a heater H_C . The Lambwave transducer LW_C is connected to inductor L_{CL} (also shown in FIG. 1). The heater H_C is connected to a resonant circuit consisting of an inductor L_{CH} and a capacitor C_{CH} . The Lamb-wave transducer LW_C acts as an agitator, mixer, or sonochemical inducer, as indicated by the connected arrows 26 in chamber 25.

A channel 27 connects the reaction chamber 25 to a detection chamber 28. The channel 27 is equipped with a Lamb-wave pump LW_{DP} , which is connected to a resonant circuit consisting of an inductor L_{DP} and a capacitor C_{DP} . The detection chamber 28 is equipped with a Lamb-wave sensor LW_D , which is connected to a capacitor C_D .

Lamb-wave transducers have high mechanical Q values and can therefore be powered by only a narrow range of alternating voltage frequencies. The Lamb-wave pumps (LW₁, LW₂, LW₃) and Lamb-wave sensor (LW_D) are powered capacitively by generating an electric field between the plates (such as plates 16 and 17 of FIG. 1 for example) at the resonant frequencies of the Lamb-wave transducers (LW₁, LW₂, LW₃, and LW_D). But, because the transducers have high Q values, only when the frequency of the imposed field is near the resonant frequency of a transducer do the transducer vibrate with any substantial magnitude. Similarly, the Lamb-wave mixing chamber transducer LW_C is provided by an alternating frequency magnetic field generated by the coil (15 in FIG. 1) at the mechanical resonant frequency of the transducer LW_C . The heater H_C and the Lamb-wave pump LW_{DP} are activated by directing an electromagnetic wave from the antenna (18 in FIG. 1) to the resonant circuit C_{CH} and L_{CH} , and resonant circuit C_{DP} and L_{DP} , respectively. The frequency of the incident electromagnetic radiation must correspond to the mechanical resonant frequency of the transducer LW_{DP} , to activate the pump LW_{DP} . The frequency of the incident electromagnetic radiation must correspond to the resonant frequency of the electrical elements C_H , L_{CH} and H_C to activate the heater \mathbf{H}_{C} .

A PCR reaction, for example, is initiated by pumping the reagents in the chamber 19, 20 and 21 along the directions of the arrows through respective channels 22, 23 and 24 to the reaction chamber 25 by activating pump LW_1 , LW_2 , and LW_3 . A series of about twenty to forty thermal cycles, for example, are then initiated, and during each cycle the temperature of the reactants in the reaction chamber 25 goes from 55° C. to 96° C., and back to 55° C., for example. The temperature of the reaction chamber 25 is determined by the power of the incident electromagnetic signal at the frequency corresponding to the resonant frequency of the circuit composed of the capacitor CC_H , and the inductor L_{CH} , together with the heater H_C . The Lamb-wave device

LW_C of the reaction chamber 25 acts as an agitator or mixer, as indicated by arrows 26, to mix the reagents and promote the reaction.

When the thermal cycling is complete, the contents of the reaction chamber 25 are pumped by the Lamb-wave perm 5 LW_{DP} through channel 27 in the direction of the arrow to the detection chamber 38, which utilizes a Lamb-wave sensor LW_D. Alternatively, the detection chamber 28 may be provided with an optical window and testing may be performed by fluorescence-based or absorption-based optical spectroscopy.

FIG. 3 illustrates a heating/detection arrangement that can be incorporated into the microfabricated reactor of FIGS. 1 and 2. As shown in FIG. 3, a chemical reaction chamber, such as a PCR chamber, of a miniaturized, microfabricated 15 instrument, generally indicated 30, is illustrated in crosssection, with chamber 31 being formed in a housing 32, constructed of Pyrex for example, and having silicon inserts 33 and 34 therein, with an inlet 35 and an outlet 36. Energy from two different energy (light) sources is directed onto the 20 housing 32, one source 37 being infrared (IR) source, and the second source 38 being an ultra-violet (UV) source. The IR source 17 applies heat more uniformly through the bulk of the solution in chamber 31. The UV source 18 induces fluorescence of the reaction products in the visible (Vis) spectrum, which can be detected by a visible (Vis) detector 39 located external of the housing 32 defining reaction chamber 31. Housing 32 must be constructed of a material transparent to UV and/or the visible spectrum. By incorporating an integrated excitation (heating) and detection system in the reaction chamber itself, confirmation of the presence of a sample in the reaction chamber can be confirmed, and the dual reaction and detection chambers 25 and 28 of the microfabricated reactor of FIG. 2 can be consolidated, thus reducing fabrication costs by reducing components.

The present invention, an embodiment of which is illustrated generally in FIGS. 4 and 5 involves a microfabricated reactor generally indicated at 40 which includes a siliconbased sleeve as a chemical reaction chamber, generally 40 indicated at 41, constructed of two bonded silicon parts, and which utilizes doped polysilicon for heating and bulk silicon for convective cooling, as described in greater detail hereinafter. The sleeve 41 includes a slot or opening 42 into which reaction fluid, indicated at 43, from a hypodermic 45 needle 44 is inserted into the reaction chamber, or into which a secondary tube 45 containing a reaction mixture 46 may be inserted. The tube 45 is constructed of plastic, for example, or other material which is inert with respect to the reaction mixture, thereby alleviating any potential material incom- 50 patibility issues. The sleeve is also provided with an opening 47 in which is located an optical window 48, made, for example, of silicon nitride, silicon dioxide, or polymers. The silicon sleeve reaction chamber 41 includes doped polysilicon for heating and bulk silicon for convective cooling, and 55 combines a critical ratio of silicon and silicon nitride to the volume of material to be heated (e.g., liquid) in order to provide uniform heating, yet low power requirements.

FIG. 6 is an enlarged view of microreaction chamber, similar to the FIG. 4 embodiment, but utilizing two windows. The reaction chamber of FIG. 6, generally indicated at 50, is composed of two silicon wafers or substrates 51 and 52 bonded together as indicated at 53, and configured to define a slot or opening 54 therein. Each of wafers 51 and 52 include a layer of silicon nitride 51' and 52' which define 65 a window, indicated generally at 55 and 56, respectively. Window 55 in wafer 51, constructed of silicon nitride, is

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provided with a heater 57 having electrical leads 58 and contacts 59 which extend along the edges of heater 57 to provide uniform heating. Window 56 in wafer 52 has a heater not shown in FIG. 6 but which is secured by metal contacts 60 and 61 as illustrated in either of FIGS. 7 and 8. The silicon nitride layers 51' and 52' are very thin (about 1 µm) and vapor-deposited onto the bulk silicon wafers 51 and 52. The silicon nitride only becomes a window, as indicated at 55 and 56, when the bulk silicon wafers 51 and 52 are etched away to form the opening or slot 54. Heater 57 is transparent to energy passing through window 55, for example.

FIG. 7 is a greatly enlarged view of an embodiment of a section of silicon wafer 52 and window 56, as indicated by the circle 62 in FIG. 6. As seen in FIG. 7, the section of the silicon wafer 52, indicated at 63, is composed of bulk or single crystal silicon and is in contact with a low (100 to 500 MPa) stress silicon nitride membrane or window 64 (52' in FIG. 6) which in turn is in contact with a doped polysilicon heater 65 and metal contact 60 and 61. The FIG. 7 embodiment comprises an isolated heater version fixed window.

FIG. 8 is a greatly enlarged view of another embodiment of a section of silicon wafer 52 and window 56, as indicated by the circle 62. As seen in FIG. 8, the sections of the silicon substate 52, indicated at 66 are composed of bulk or single crystal silicon. As in the FIG. 7 embodiment, a low (100 to 500 MPa) stress silicon nitride member or window 69 (52' in FIG. 6) is in contact with silicon section 66, a doped polysilicon heater 70 is in contact with window membrane 69 and metal contacts 71 are mounted to heater 70. The FIG. 8 embodiment comprises a non-isolated heater version. The window size relative to the chamber can be varied to ensure thermal uniformity and optical access to the reaction chamber.

By way of example, the silicon wafers or substrates 51 and 52 may have a length of 5 to 50 mm, width of 2 to 10 mm, thickness of 0.1 to 1.0 mm, with the slot 54 having a cross-sectional area of 5 to 500 mm². Slot 54, which shown to be of a six-sided configuration, may be a round, oblong, square, rectangular, or other configuration. Windows 55 and 56 may have a length of 0.1 to 1 mm, width of 0.1 to 50 mm, thickness of 0.1 to 10 µm, and in addition to silicon nitride, may be composed of silicon dioxide, silicon, or polymers. The doped polysilicon heater 65 of FIG. 7 may have a thickness of 0.05 to 5 μm , with the heater 70 of FIG. 8 having a thickness of 0.05 to 5 µm. The metal contacts 60-61 and 61' of FIGS. 6 and 7 may be composed of gold or aluminum, with a thickness of 0.01 to 5 µm, with the metal contact 71 of FIG. having a thickness of 0.01 to 5 µm and composed of gold or aluminum. The heater 57 in silicon wafer or substrate 51 is composed of doped polysilicon having a thickness of 0.05 to $5 \mu m$, with the electrical leads and contacts 58 and 59 being composed of gold or aluminum.

The use of bulk silicon, polysilicon, silicon nitride enables flexibility in design for thermal and optical properties of each chamber. This enables individually controlled, thermally isolated reaction chambers in a small instrument (FIG. 9) or in large instrument (FIG. 10).

FIG. 9 is an embodiment of a miniature thermal cycling, battery operated, hand-held low-power, feedback-controlled instrument for PCR that uses microfabricated, silicon-based reaction chambers, such as those of FIGS. 4 and 6, the development of which addressed thermal uniformity and temperature precision of the reaction chambers, temperature ramp rates of the chambers, and biocompatibility of the materials in contact with the reagents.

As shown in FIG. 9, the hand-held, battery-operated instrument, coined "PCR man", generally indicated at 75, comprises a pressure-fit electrical contact controller holder, or housing 76, which for example may be 3×5 inches having a control-face-plate 77 with various indicators thereon, 5 including a "status" window 78. The holder 76 is provided with a thermocouple-based temperature feedback control circuitry, heater electronics, computer interface, and power source connector, as described in greater detail hereinafter. The holder 76 is provided with batteries, indicated at 79, 10 such as four nine-volt batteries, and at the upper end is provided with slots 80 for insertion of reaction chambers inside the holder (three slots shown), and into which one or more silicon-based reaction chambers 81, 82, 83 and 84, with integrated heaters (as shown in FIG. 6) are inserted as indicated by the arrow 85. The reaction chambers 81–84 ¹⁵ may when constructed contain different reagents or chemicals, and can be selectively inserted into the handheld instrument 75 via slots 80 in holder or controller 76.

This instrument can be used to rapidly and repetitively provide controlled thermal cycles to the reaction mixture. The thermal conductivity properties of the silicon or similar semiconducting substrate, help speed up the thermal rise and fall times, and allow low power operation. While silicon is unique in its thermal properties, i.e., high thermal conductivity, a combination of silicon, silicon nitride, silicon dioxide, polymers and other materials would provide a combination of thermal conductivity and insulation that would allow thermal uniformity and low power operation.

The particular embodiment, such as FIG. 6, of a microfabricated reactor described can be used as a thermal cycling instrumentation for use inthe PCR and other chemical reactions, biochemical processes, microbiological processes, and incubators. As shown hereinafter the reaction chamber of this invention is superior to present commercial instruments used in thermally-driven chemical reactions.

During the experimental verification of the instrument of FIG. 9 and the microreaction chambers for use therein, such as illustrated in FIGS. 4 and 6, several different sizes of PCR reaction chamber designs were fabricated using integrated 40 circuit (IC)-type silicon processing steps. The generalized fabrication process was as follows: Three-inch round, 0.5 mm thick single crystal silicon (SCS) wafers were processed inthe following way: low stress (200–300 MPa) silicon nitride (Si_xN_v) was low-pressure chemical vapor (LPCVD) 45 deposited onto entire wafer (1.0–2.0 µm thick). Photolithographic patterns for reaction chamber and subsequent processing steps were taken in the following order: 1) the silicon nitride was reactive ion etched (RIE) over the reaction chamber area, 2) the SCS was etched to the silicon 50 nitride backside defining the chamber volume, 3) the wafer was patterned and the silicon nitride is chemically etched away everywhere except over the nitride membrane or left over the entire surface, depending upon the reaction chamber design, 4) the remaining silicon nitride membrane (side 55 opposite the chamber) was LPCVD deposited with polycrystalline silicon (polysilicon) to a thickness of 3000Å, 5) the polysilicon was then high temperature doped with boron to a resistivity of 50–200 ohms per square, and 6) either aluminum or gold thin-film metal contacts were deposited 60 defining the heater geometry.

Each wafer potentially contains many reaction chambers, depending upon geometry and volume desired. The etched depression in each wafer constitutes one-half of a dual-heater reaction chamber. Processed wafers are subsequently 65 bound together forming an enclosed chamber with heaters on both sides.

The reaction chambers can be bonded together by depositing a thin film of low-temperature-curing polyimide between the two wafers directly or other bonding techniques such as eutectic metal bonding. A high precision computer-controlled silicon saw was used in each design to cut out each dual-heater chamber. The chambers were then rinsed repeatedly with de-ionized water and dried prior to treatment with silane.

The reaction chambers were inserted into a pressure-fit electrical contact holder that was part of the plexiglas backboard of the electronics components making up the controller. The controller electronics could be either/or anologue or digital and could use processes such as pulse-width modulation as a feedback control mechanism. The backboard was 3 inches by 5 inches and consisted of the thermocouple-based temperature feedback control circuitry, heater electronics, computer interface, and power source connector. The circuitry was designed to work from 8 to 32 volts. Thermal calibration was accomplished by correlating the temperature of the fluid with that of the silicon-measuring Type K thermocouple. Once calibrated, the instrument was capable of automated, feedback-controlled, thermal cycling operation without direct measurement of the reaction fluid. The thermal cycler output is to an Apple Centris 650 computer which displays the thermal cycle real-time along with storing the accumulated profiles. Four nine-volt batteries were abel to run the entire instrument continuously for over 2.5 hours.

Typical PCRs were set up as scaled-up master mixes, to assure uniformity between aliquotes thermocycled under different conditions. Reagent amounts were based on those ideal for 50 ul reactions. In general, master mixes contained: 50 Mm KCl, 10 mM Tris-HCl pH 8.3, 1.5-3.0 mM MgCl₂, 200 uM each deoxynucleotide, or 800 uM dNTP total, 0.5 uM each of two oligonucleotide primers, 25 units/ml Ampli-Taq® DNA polymerase, and target template at a specified copy number per 50 ul reaction. Template for some of the β-globin PCRs was added as single strand DNA from a M13 bacteriophage clone of a portion of the human β -globin gene. CF template was human genomic, double stranded, DNA derived from a cultured cell lines, HL60, GM07460, or GM08345. Each reaction mixture was aliquoted from the same master mix and thermocycled in the instrument of the present invention and a Perkin-Elmer GeneAmp® 9600 Thermal Cycler. Thermocycled reactions from both thermal cyclers were fractionated on 3% NuSeive, 1% Seakem agarose (FMC Corp.) using tris-borate buffer. The gels were stained with ethidium bromide and photographed under illumination with 302 nm UV light.

Although initially conceived as single use, disposable reaction chamber, the robust nature and stable properties allowed for repeated use of the reaction chambers.

The (MEMS) based thermal cycling instrument of this invention has been tested with a variety of PCR systems, including viral, bacterial, and human genomic templates. As well, various changes in both the reaction chamber design and controller instrumentation have been implemented and evaluated. A controller output real-time display of a thermal cycle from microfabricated thermal cycler has been prepared and it has been shown that with 15 volts input (average 1.2 Watts) that heating rates of over 5° C./sec are attained. Cooling is slightly slower (2.5° C./sec.) mostly due to the fact that the reaction chamber is held inside a plexiglass instrument board. Precision of ±0.5° C. is maintained at the target temperatures. Higher heating and cooling rates have been achieved.

We have performed experiments that show the quantitative nature of the PCR process in both FIG. 9 and commer-

cial instruments. These experiments consisted of removing 5 μ L aliquots out of a **105** starting copies, β -globin PCR from both the instruments at 23, 25, 27, 29, and 31 cycles. These aliquots were subsequently run on an agarose electrophoresis gel. The results from both instruments are virtually identical. The same quantitative gel electrophoresis series results from the amplification of the 268-bp target of β -globin directly from human genomic (HL60) DNA were per

Multiplex PCR is considered to one of the most recent and 10 analytically-powerful DNA amplification techniques. It requires precise and uniform temperature control within the reaction chamber. We have achieved this with the instrument of this invention.

Post-PCR-detection of the specific mutations associated with the cystic fibrosis (CF) disease, for example, can be identified with simple nylon-based test strips, using reverse-dot-blot technology. The test strip has specific, immobilized DNA probes containing the mutation sequence of interest. The multiplex PCR amplification products are put into a simple reagent trough along with the assay. If binding occurs and the DNA is retained after a wash step, the DNA-biotin-streptavidin-enzyme complex will turn color upon treatment with the substrate. The commercial and the FIG. 9 instrument-amplified results of PCR followed by reverse-dot-plot assay for CF prepared.

From the results of the above-referenced experiments and previous results, relative to the above-identified copending applications, with single-sided heaters, silicon-based reaction chambers of various sizes and configurations are capable of carrying out chemical reactions, such as PCR, with low power requirements.

The significance of the above-reference experimental results is that for the first time, battery-operated, hand-held, PCR amplification; and simple reagent-based, targeted detection of complex biologicals and diseases can be carried out in an instrument such as illustrated in FIG. 9.

The rapid temperature cycling and thermal uniformity now possible in a PCR-type compatible silicon-based 40 microreaction chamber may provide insight into hybridization and enzyme kinetics. For example, the importance of temperature control is paramount in the PCP process, especially when complex systems are to be amplified (e.g., human genomic DNA, multiplex amplifications). Precise 45 temperature control as well as thermal uniformity must be balanced. To truly miniaturize the instrument or take advantage of microfabricated reaction chambers in order to build a high-throughput instrumentation, such as illustrated in FIGS. 10A, 10B and 11, one must integrate the control 50 elements on a unit-by-unit scale. Thermal properties of the various materials used must also be balanced to combine efficient control with thermal liability. Silicon-based materials afford the requisite thermal properties, the ability to integrate heaters and feedback control, and their manufac- 55 ture takes advantage of highly parallel, automated, and batched processing.

FIGS. 10A-10B and 11 illustrate a system approach, combining the high-throughput, high efficiency thermal cycler instrument, sample handling, and electrophoresis 60 modul. The electrophoresis module could also be micromachined in glass or silicon. The instrument could be hybrid in nature; i.e., a silicon based reaction chamber and a mini glass electrophoresis module taking advantage of both substrates or members, as in the FIG. 5 embodiment. The 65 advantage to having real-time detection of DNA production is that it allows the operator to know about the PCR

efficiency during the reaction, rather than waiting to see the results on a gel. This will significantly help DNA sequencing productivity by eliminating time wasted running electrophoresis gels on samples that haven't amplified.

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FIGS. 10A and 10B illustrate a thermal cycling instrument, generally indicated at 90, having a housing 91 with a face plate 92 with various indicators thereon, including a "status" window 93, similar to the faceplate of the FIG. 9 hand-held instrument. The housing includes a hinged top 94, under which is located an array 95 (see FIG. 10B) of individually controlled silicon-based microreaction chambers 96, which may, for example, be of the type illustrated in FIGS. 4 and 6. The instrument 90 is designed for 384 microreaction chambers 95, although the array 95 as shown in FIG. 10B only includes 100 chambers for simplicity of illustration.

FIG. 11 is a schematic representation of high-throughput DNA application, sample-handling, and electrosystem utilizing the instrument of FIGS. 10A-10B, and corresponding reference numeral indicate corresponding components. An array 95' of 384 individual-controlled PCR reaction chambers 96' (only five shown, is operatively connected to an automated sample input/output assembly, generally indicated at 97 using two sets of microinjectors, generally indicated at 98 and 99. The sample input/output function between microinjector set 98 of assembly 97 and array 95 is indicated by double arrow 100, while the function between the sets 98 and 99 of microinjectors is indicated by double arrow 101. The microinjector set 99 is operatively to an array 102 of individual microelectrophoresis channels 103. This injector input/output system will load reagent samples from the reaction chambers 96 with vacuum or electrokinetic power; automatically or robotically move to electrophoresis channels 103; and unload reagents via pressure or reversed field electrokinetic injection into those channels for electrophoretic separation. The electrophoresis module could be micromachined as well. Silicon is good for reaction chambers, glass for electrophoresis.

The electrophoresis channels 103, formed in a glass substrate are each directly connected to a silicon reaction chamber of the type shown in FIG. 4, so as to produce an array 95 of reaction chambers 96' connected directly to the array 102 of electrophoresis channels 103, as shown in FIG. 5.

Removable/permanent lines/inserts for the reaction chambers of a material kown to be compatible with the appropriate reactions, such as shown in FIG. 12 will in some applications reduce the overall cost, as these liners/inserts may be disposable. Also, considered are derivatizing agents for the surfaces of the silicon-based reaction chamber to enhance covalent and/or other bonding to the liners. Examples being the organic/reactive silanes, polyimides, teflons, polytheylene, other polymers.

FIG. 12 illustrates an embodiment of an insert/liner, generally indicated at 105, for a reaction chamber with an optical window 106 therein. The insert/liner 105 includes a six-sided housing 107 and a top/cover 108. The six-sided housing 107 is configured, for exmple, to be inserted into opening 54 of the reaction chamber 50 of the FIG. 6 embodiment, such that window 106 aligns with one of windows 55 or 56 of FIG. 6. The housing 107 may be constructed of plastic or other compatible material set forth above. Window 106 of insert/liner 105 includes a test strip 109, described hereinafter with respect to FIG. 14.

FIG. 13 illustrates external filling of the reaction chamber insert/liner 105 of FIG. 12 via an external interfludic con-

nection, generally indicated at 110. Examples of fluidic connections includes: syringe needles, pipette tips, and fused silica capillaries or glass or polymer tubing.

Surface immobilization of the windows (or test strip) with probes for optical or other detection (other microbased detections) of product production and specificity, can be provided as shown in FIG. 14 which is an enlarged view of the test strip 109 of FIG. 12. Such a test strip can be included in the windows of the FIGS. 4 or 6 reaction chambers. Immobilized reagents/probes for detection of specific products directly on the window, such as 106 of FIG. 12, or within the reaction fluid in reaction chamber insert/liner 105 of FIG. 12, can be detected optically inthe PCR man hand-held instrument of FIG. 9, by the use of the test strip 109. The actual inner surface of the window could be used 15 as an immobilization surface for specific-target or product detecting probes, or the window could be used to view an immobilization/detection surface within the chamber.

FIGS. 15 and 16 schematically illustrate two setups for optical detection. The FIG. 15 setup is a laser/ccd version, while the FIG. 16 setup will allow low-power operation for implementation into the PCR man (hand-held instrument) of FIG. 9.

As shown in FIG. 15, this optical detection arrangement for a reaction chamber 120 with a window 121 and control electronics 122, includes an optical filter 123, such as an interference filter or band pass filter for passing the detection wavelength of interest, CCD 124, digitized image generally indicated at 125, focusing optics 126, reflector/splitter 127 and an Argon ion laser 128. The operation is as follows: The laser excites the fluorescent indicator dye associated with product detection. The fluorescent signal is monitored by the CCD 124. Absorption spectroscopy could similarly be used.

FIG. 16 is a miniaturized optical detector system for reaction chamber 120' having a window 121' and control electronics 122' is composed of two filters 130 and 131, a solid state detector 132 and a Blue LED 133. The filters 130 and 131 are either band pass or long pass for selecting emission (i.e., 600 nm long pass) and band pass for selecting the excitation wavelength of interest, such as 488 nm±10 nm. The excitation band pass can be used to select from the typically broad emission of an LED, for example. The operation of the FIG. 16 detection system is as follows: The LED is filtered to 488+10 nm as an excitation source (or absorption) for the fluorescent indicating dye. The solid state detector is also filtered to receive only the wavelengths of detection (>600 nm) or as an absorption detector.

Artificial intelligence is one way to produce DNA and determine how many cycles to go, when it is complete, if it 50 worked, adjustment of parameters to improve production, etc. Using a real-time detection systems such as illustrated schematically in FIG. 17, an artificial intelligent feedback system using integrated detection can be provided. The system of FIG. 17 comprises a reaction chamber 135 having 55 a window 136, a detector 137 for in situ detection of DNA production, an instrument control 138 for reaction chamber 135, and a data readout system 139, which receives data from detector 137, as indicated by arrow 140, and supplies control data to controller 138, as indicated by arrow 141. 60 The data readout system 139 provides information such as how much DNA is being made, starting copy number, reaction complete, etc. By quantifying the DNA production via the optical monitoring system, which is well known, the system could adjust its cycling time and cycle number to 65 produce the minimal number of cycles required for detection, thus speeding up the process. Also by determining the

cycle number required to detect a given fluorescent signal, or product concentration, the system would be able to calculate all starting copy number or concentration of the unknown starting sample. This would allow automated concentration calculations. Real-time quantitative information can allow the system to adjust the reaction parameters such as target temperatures, hold times, and ramp rates.

A microfabricated, electrochemiluminesence cell for the detection of amplified DNA is described hereinafter with respect to FIGS. 18–31, and which sets forth the design, fabrication, and testing thereof. The microcell is designed to be the detection unit in a PCR micro-instrument, such as described above and illustrated in FIG. 9. The cell is a vertical assembly of micromachined silicon and glass and contains thin film electrodes, as shown in the Figures.

The detection of DNA by means of electrochemiluminescence starts with DNA amplification by PCR, to increase the concentration to detectable levels. Then it is labeled with tris (2,2' bipyridyl) ruthenium (II) (TBR). Oxidized TBR luminesces (orange) upon reduction. Oxidation occurs electrochemically at an electrode surface, hence the light emission is referred to as electrochemiluminescence (ECL). TBR requires a relatively low oxidation potential (a few volts) and has a high ECL efficiency in the visible (620 nm). This makes it attractive for microsensor applications, since visible emission is readily detected with silicon photodiodes, which could be integrated into a silicon micromachined cell. The reduction can occur electrochemically or chemically; in either case, light is emitted. For example, oxidized tripropylamine (TPA) readily transfers an electron to oxidized TBR, whereupon the TBR chemiluminesces. Since both oxidations can occur at the same electrode, relatively large concentrations of both species can be produced in close proximity, which results in higher light intensity for a given TBR concentration, than if TBR alone is present in solution. The electrochemical oxidation and chemical reduction reactions for TBR which occurs at the anode are schematically diagrammed in FIG. 18. Electrochemical reduction of TBR also occurs at the cathode. In order to oxidize only the TBR labeled DNA and not the free TBR, a separation of the two is required. One way to achieve this is by using the highly specific binding of immunoproteins (antibody-antigen).

An example is shown in FIG. 19, where a biotin primer is made on a 5' end of one strand of target DNA and the TBR is tagged to the 5' end of the complementary strand. During the PCR process DNA double strands are produced with biotin and TBR labeled on either end. The biotin labeled DNA can then be introduced into an electrochemical cell with an anode whose surface is coated with avidin, the antibody for bitoin. Selective binding will occur, after which the solution in the cell is flushed to remove any "free" TBR. Now the TBR, bound to the DNA, which in turn is attached to the anode via the antibody-antigen bond, can be oxidized along with added TPA, and the subsequent luminescence intensity will depend on the amount of DNA that is present.

The ECL microcell, as described in greater detail hereinafter with respect to FIGS. 21–31, is a multilayer assembly of micromachined silicon and glass. Cells with solution capacity ranging from 35 µL to 85 µL have been designed and fabricated in silicon. An e-beam deposited, gold, thin film forms the cell cathode. The anode is also a thin film. Experiments with both indium tin oxide (ITO) and platinum have been carried out. ITO is transparent to visible light, so that when deposited onto glass, it can form the top layer of the assembly, through which the emitted light can be picked up by a photodetector (see FIG. 21). The assembly also contains micromachined fluid fill ports (see FIG. 22). The

layers were assembled and bonded together (see FIGS. 29–30) using a low temperature curing polyimide, such as Epotek 400.

ECL experiments have been performed in the microcell with free TBR, i.e., no DNA. The cells were filled with 5 TPA+TBR solution and a photomultiplier tube (PMT) was placed in close proximity to the top glass layer of the cell to detect emission. The chemiluminescence produced by the reaction of oxidized TPA and TBR depends on the concentration of both chemicals. In these experiments, the concentration of TPA was kept constant (50 mM) and TBR was varied. The solutions were prepared as follows: 1 g of TBR hexahydrate chloride was dissolved in 50 mM TPA to make 5 mM of TBR, which was then diluted with additional 50 mM TPA to produce a set of test solutions, whose TBR concentrations range from 0.1 nM to 5 mM. An EG&G potentiostat, model PARC 273, was used to produce voltammograms of the TBR+TPA solution, both in the microcell with ITO and gold thin film electrodes, and in a more conventional, electrochemical cell with platinum wire electrodes. From the voltammogram, the oxidation potential, ²⁰ which is where ECL occurs, was determined and then applied as a dc bias between the thin film cathode and anode. The emitted light was measured with a Hamamatsu MT, model R928, biased at 600 volt. FIG. 20 shows the relationship between measured light intensity and electrode 25 voltage for a TBR concentration of /mM, where cell voltage and ECL intensity versus time. The voltage, as indicated by the dot-dash-dot line, is increased, then decreased. In both directions, the voltage passes through the oxidation potential of TBR, where intensity of ECL is a maximum. In tests 30 conducted thus far, the lowest concentration of TBR that has been measured Using the microcell with an ITO film as the anode material was 1 µM. With a platinum anode, the measured TBP concentrations were as low as 1 nM. The relatively high resistance of the ITO film is believed to be limiting the oxidation current for TPA, and therefore reducing the sensitivity. It has been determined that sensitivity can be improved by depositing a thin film of material, such as aluminum on the ITO film, as described hereinafter with respect to FIG. 31. Also, efforts are being carried out to integrate the silicon photodiode into the microcell, rather 40 than being separated therefrom as in the FIG. 21 embodiment.

FIG. 21 illustrates an embodiment of a micromachined ECL cell with thin film anode, generally indicated at 140, and a silicon (Si) photodiode 141 positioned adjacent the 45 ECL cell 140. The ECL cell 140 is shown in enlarged cross-section in FIG. 22. The cell 140 comprises a pair of silicon members 142 and 143, between which is positioned an electrode 144, which may be constructed of gold (Au), platinum (Pt) or silver (Ag), an ITO layer 145, and a glass 50 layer or slide 146. Silicon member 142 includes a reaction chamber 147, and member 143 includes a pair of filling ports 148 (see FIG. 22) via which an analyte, as indicated by legend is directed into chamber 147 and withdrawn therefrom via tubes or lines 149 and 150, as indicated by arrows 55 151 and 152. As seen in FIG. 22, a center section 153 of silicon member 143 located between fill ports 148, along with ITO layer 145 and glass slide 146 define a window by which reactions within chamber 147 can be detected, as indicated by photons 154 passing therethrough onto photo- 60 diode 141. Electrical leads 155 and 156 are connected from a power source to electrode 144 and ITO layer 145, respectively, while photodiode 141 is electrically connected to a power source via leads 157 and 158.

FIGS. 23–30 illustrate the fabrication of an embodiment 65 of an ECL cell similar to that of FIGS. 21 and 22. The fabrication process is carried out as follows:

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- 1. A block 160 of silicon is coated to form a layer 161 of silicon nitride (see FIG. 23).
- 2. A layer 162 of photoresist is deposited on the layer 161 (see FIG. 24).
- 3. The layer 162 is patterned and photolithographic process to form an opening 163 therein (see FIG. 25).
- 4. The section 161' of silicon nitride layer 161 beneath the opening 163 is removed by RIE etching (see FIG. 26).
- 5. A section of silicon block 160 is removed by KOH etching to form a reaction chamber 164, and the remaining photoresist 162 is removed (see FIG. 27).
- 6. A layer of gold, for example, is deposited by thin film evaporation over the upper surface of block 160 and chamber 164 to form an electrode 165 (see FIG. 28).
- 7. A second block of silicon 166 is coated with a layer 167 of silicon nitride and openings 168 and 169 are formed therein by RIE etching, and a pair of filling ports 170 and 171 are formed, as by micromachining, in block, 166, and silicon nitride coated block 166 is bonded to electrode 165 (see FIG. 29).
- 8. A layer of ITO forming an electrode 172 is deposited on a layer or slide 173 of glass, and then bonded to the silicon nitride layer 167 (see FIG. 29).
- 9. Electrical leads 174 and 175 are secured to gold electrode 165 and ITO electrode 172, a detector 176, such as the photodiode of FIG. 21, having electrical leads 177 and 178 is bonded to glass layer 173, and the silicon nitride coated silicon block 160 is positioned on a magnet 179 having electrical leads 180 and 181 (see FIG. 30).

To reduce resistance of the ITO electrode 172 a thin film of aluminum 182 (see FIG. 31) can be deposited on the ITO layer or electrode 172 prior to same being bonded to the silicon nitride coated silicon block 166.

It has thus been shown that the present invention provides a silicon-based microreaction chamber which can be used in a hand-held instrument or a large high-throughput instrument. In addition, the invention provides for insert/liners, test strips, optical detection, and automatic control for the microreaction chamber. Thus, the present invention substantially advances the state of the art for PCR and other chemical reactions.

While particular embodiments, materials, parameters, etc. have been set forth to exemplify and explain the principles of the invention, such are not intended to be limiting. Modifications and changes may become apparent to those skilled in the art, and it is intended that the invention be limited only by the scope of the appended claims.

We claim:

- 1. In a microfabricated chemical reactor having a reaction chamber, the improvement comprising:
 - a sleeve reaction chamber,
 - said sleeve reaction chamber having a slot therein,
 - said slot being constructed to enable insertion of an insert or liner therein, and

heating means for said sleeve reaction chamber.

- 2. The improvement of claim 1, wherein said slot is so constructed as to enable insertion of reaction fluid therein either directly or via a tube.
- 3. The improvement of claim 1, wherein said sleeve reaction chamber is provided with at least one optical window.
- 4. The improvement of claim 1, wherein said sleeve reaction chamber is composed of a plurality of bonded silicon members.

- 5. The improvement of claim 4, wherein said bonded silicon members are constructed of polysilicon and bulk silicon.
- 6. The improvement of claim 5, wherein doped polysilicon constitutes said heating means and bulk silicon is a 5 convective cooling means.
- 7. The improvement of claim 1, wherein said sleeve reaction chamber includes a pair of windows and wherein said heating means is located adjacent each of said windows.
- 8. The improvement of claim 7, wherein said windows are 10 constructed of silicon nitride.
- 9. The improvement of claim 7, wherein said heating means comprises a doped polysilicon heater.
- 10. The improvement of claim 1, wherein said slot in said reaction chamber is of a multi-sided configuration.
- 11. The microfabricated chemical reactor of claim 1, wherein said sleeve reaction chamber is provided with at least one window.
- 12. The microfabricated chemical reactor of claim 11, additionally including an insert adapted to be inserted into 20 said slot, said insert including at least one window.
- 13. The microfabricated chemical reactor of claim 12, wherein said window of at least said insert is provided with a test strip.
- 14. The improvement of claim 1, wherein said silicon- 25 based sleeve reaction chamber is in combination with and constructed to be inserted into a hand-held, battery-operated instrument.

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- 15. The improvement of claim 1, wherein said silicon-based sleeve reaction chamber is in combination with and constructed to be inserted into an instrument constructed to contain an array of such reaction chamber.
- 16. The improvement of claim 15, wherein said array of reaction chambers is operatively connected via an array of microinjectors to a microelectrophoresis array.
- 17. The improvement of claim 15, wherein said array of reaction chambers is connected directly to the microelectrophoresis array.
- 18. The improvement of claim 17, wherein said array of reaction chambers is constructed of silicon and wherein said microelectrophoresis array is constructed of glass.
- 19. The improvement of claim 3, additionally including an optical detector positioned adjacent said optical window.
- 20. The improvement of claim 19, additionally including a data readout system operatively connected to said optical detector, and an instrument controller operatively connected to said data readout system and said reaction chamber.
- 21. The improvement of claim 1, additionally including a liner adapted to be inserted into said slot.
- 22. The improvement of claim 1, wherein said reaction chamber is silicon-based and provided with at least one window adjacent said slot, and wherein said heating means comprises a heater positioned adjacent said window.

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